

FORM PT-100 (Modified)
(REV 11-2000)

U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE

ATTORNEY'S DOCKET NUMBER

TRANSMITTAL LETTER TO THE UNITED STATES

SHIG C11119

DESIGNATED/ELECTED OFFICE (DO/EO/US)

U.S. APPLICATION NO. (IF KNOWN, SEE 37 CFR

CONCERNING A FILING UNDER 35 U.S.C. 371

09/913625

INTERNATIONAL APPLICATION NO.

PCT/JP00/02056

INTERNATIONAL FILING DATE

30.03.00 (30 March 2000)

PRIORITY DATE CLAIMED

17.12.99 (17 December 1999)

TITLE OF INVENTION

HYDROGEN STORAGE METAL ALLOY AND PRODUCTION THEREOF

APPLICANT(S) FOR DO/EO/US

OKADA, Masuo and KURIIWA, Takahiro

Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:

1. ☒ This is a **FIRST** submission of items concerning a filing under 35 U.S.C. 371.
2. ☐ This is a **SECOND** or **SUBSEQUENT** submission of items concerning a filing under 35 U.S.C. 371.
3. ☒ This is an express request to begin national examination procedures (35 U.S.C. 371(f)). The submission must include items (5), (6), (9) and (24) indicated below.
4. ☐ The US has been elected by the expiration of 19 months from the priority date (Article 31).
5. ☒ A copy of the International Application as filed (35 U.S.C. 371 (c) (2))
 - a. ☒ is attached hereto (required only if not communicated by the International Bureau).
 - b. ☐ has been communicated by the International Bureau.
 - c. ☐ is not required, as the application was filed in the United States Receiving Office (RO/US).
6. ☒ An English language translation of the International Application as filed (35 U.S.C. 371(c)(2)).
 - a. ☒ is attached hereto.
 - b. ☐ has been previously submitted under 35 U.S.C. 154(d)(4).
7. ☒ Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371 (c)(3))
 - a. ☐ are attached hereto (required only if not communicated by the International Bureau).
 - b. ☐ have been communicated by the International Bureau.
 - c. ☐ have not been made; however, the time limit for making such amendments has NOT expired.
 - d. ☒ have not been made and will not be made.
8. ☐ An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).
9. ☒ An oath or declaration of the inventor(s) (35 U.S.C. 371 (c)(4)).
10. ☐ An English language translation of the annexes of the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371 (c)(5)).
11. ☐ A copy of the International Preliminary Examination Report (PCT/IPEA/409).
12. ☒ A copy of the International Search Report (PCT/ISA/210).

Items 13 to 20 below concern document(s) or information included:

13. ☐ An Information Disclosure Statement under 37 CFR 1.97 and 1.98.
14. ☒ An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.
15. ☒ A **FIRST** preliminary amendment.
16. ☐ A **SECOND** or **SUBSEQUENT** preliminary amendment.
17. ☐ A substitute specification.
18. ☐ A change of power of attorney and/or address letter.
19. ☐ A computer-readable form of the sequence listing in accordance with PCT Rule 13ter.2 and 35 U.S.C. 1.821 - 1.825.
20. ☐ A second copy of the published international application under 35 U.S.C. 154(d)(4).
21. ☐ A second copy of the English language translation of the international application under 35 U.S.C. 154(d)(4).
22. ☒ Certificate of Mailing by Express Mail
23. ☐ Other items or information:

U.S. APPLICATION NO. (IF KNOWN, SEE 37 CFR

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INTERNATIONAL APPLICATION NO.

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24. The following fees are submitted:

CALCULATIONS PTO USE ONLY

BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5)) :

- ☐ Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(a)(2)) paid to USPTO and International Search Report not prepared by the EPO or JPO \$1000.00
- ☒ International preliminary examination fee (37 CFR 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO \$860.00
- ☐ International preliminary examination fee (37 CFR 1.482) not paid to USPTO but international search fee (37 CFR 1.445(a)(2)) paid to USPTO \$710.00
- ☐ International preliminary examination fee (37 CFR 1.482) paid to USPTO but all claims did not satisfy provisions of PCT Article 33(1)-(4) \$690.00
- ☐ International preliminary examination fee (37 CFR 1.482) paid to USPTO and all claims satisfied provisions of PCT Article 33(1)-(4) \$100.00

ENTER APPROPRIATE BASIC FEE AMOUNT =

\$860.00

Surcharge of \$130.00 for furnishing the oath or declaration later than ☐ 20 ☐ 30 months from the earliest claimed priority date (37 CFR 1.492 (e)).

\$0.00

CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE
Total claims	21 - 20 =	1	x \$18.00
Independent claims	4 - 3 =	1	x \$80.00

\$18.00

\$80.00

Multiple Dependent Claims (check if applicable).

☐

\$0.00

TOTAL OF ABOVE CALCULATIONS =

\$958.00

- ☒ Applicant claims small entity status. (See 37 CFR 1.27). The fees indicated above are reduced by 1/2.

\$479.00

SUBTOTAL =

\$479.00

Processing fee of \$130.00 for furnishing the English translation later than ☐ 20 ☐ 30 months from the earliest claimed priority date (37 CFR 1.492 (f)).

\$0.00

TOTAL NATIONAL FEE =

\$479.00

Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31) (check if applicable).

☒

\$40.00

TOTAL FEES ENCLOSED =

\$519.00

Amount to be:
refunded

\$

charged

\$

- a. ☐ A check in the amount of _____ to cover the above fees is enclosed.
- b. ☐ Please charge my Deposit Account No. _____ in the amount of _____ to cover the above fees. A duplicate copy of this sheet is enclosed.
- c. ☒ The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 08-1391 A duplicate copy of this sheet is enclosed.
- d. ☒ Fees are to be charged to a credit card. **WARNING:** Information on this form may become public. **Credit card information should not be included on this form.** Provide credit card information and authorization on PTO-2038.

NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.

SEND ALL CORRESPONDENCE TO:

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24,315

REGISTRATION NUMBER

August 16, 2001

DATE

09/913625

518 Rec'd PCT/PTO 16 AUG 2001

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Appln. Of: OKADA et al.
For: Hydrogen Storage Metal Alloy and Production Thereof
DOCKET: SHIG C 11119

Assistant Commissioner of Patents
Washington, D.C. 20231

PRELIMINARY AMENDMENT

Dear Sir:

Applicant respectfully requests that the following amendments be made to the above-identified application prior to examination.

IN THE CLAIMS:

Please amend claims 6, 8, 11 and 12 to read as follows:

6. (Amended) The hydrogen storage metal alloy according to Claim 1 wherein the element X is admixed at its atom % concentration, d (at%), ranging within $0 \leq d \text{ (at\%)} \leq 20$, the atomic radius of which is larger than that of Cr but smaller than that of Ti.

8. (Amended) The hydrogen storage metal alloy according to Claim 1 wherein the element T is admixed at its atom % concentration, e (at%), ranging within $0 \leq e \text{ (at\%)} \leq 10$ and includes at least one or more members selected from the group

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consisting of Nb, Ta, Mn, Fe, Al, B, C, Co, Cu, Ga, Ge, Ln (a variety of lanthanoid metals), N, Ni, P, and Si.

11. (Amended) The process of Claim 9 wherein the predetermined time range at the aforementioned heat treatment is from 1 minutes to 100 hours.

12. (Amended) The process according to Claim 9 wherein the element ratios are those described in Claim 1 regarding the hydrogen storage metal alloy products.

Please add new claims 13-21 reading as follows:

--13. The hydrogen storage metal alloy according to Claim 3 wherein the element X is admixed at its atom % concentration, d (at %), ranging within $0 \leq d \text{ (at \%)} \leq 20$, the atomic radius of which is larger than that of Cr but smaller than that of Ti.

14. The hydrogen storage metal alloy according to Claim 13 wherein the element X is at least one or more members selected from the group consisting of Al, Ge, Ga, Si, Au and Pt.

15. The hydrogen storage metal alloy according to Claim 3 wherein the element T is admixed at its atom % concentration, e (at%), ranging within $0 \leq e \text{ (at\%)} \leq 10$ and includes at least one or more members selected from the group consisting of Nb, Ta, Mn, Fe, Al, B, C, Co, Cu, Ga, Ge, Ln (a variety of lanthanoid metals), N, Ni, P, and Si.

16. The hydrogen storage metal alloy according to Claim 5 wherein the element X is admixed at its atom % concentration, d (at%), ranging within $0 \leq d \text{ (at\%)} \leq 20$, the atomic radius of which is larger than that of Cr but smaller than that of Ti.

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17. The hydrogen storage metal alloy according to Claim 16 wherein the element X is at least one or more members selected from the group consisting of Al, Ge, Ga, Si, Au and Pt.

18. The hydrogen storage metal alloy according to Claim 5 wherein the element T is admixed at its atom % concentration, e (at%), ranging within $0 \leq e \text{ (at\%)} \leq 10$ and includes at least one or more members selected from the group consisting of Nb, Ta, Mn, Fe, Al, B, C, Co, Cu, Ga, Ge, Ln (a variety of lanthanoid metals), N, Ni, P, and Si.

19. The process of Claim 10 wherein the predetermined time range at the aforementioned heat treatment is from 1 minute to 100 hours.

20. The process according Claim 9 wherein the element ratios are those described in Claim 3 regarding the hydrogen storage metal alloy products.

21. The process according to Claim 9 wherein the element ratios are those described in claim 5 regarding the hydrogen storage metal alloy products.--

REMARKS

The claims have been amended to remove multiple dependencies. New claims 13-21 have been added to further scope the claims. Applicant requests that the filing fee be calculated based on the claims as amended.

In the event there are any fee deficiencies or additional fees are payable, please

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charge them (or credit any overpayment) to our Deposit Account No. 08-1391.

Respectfully submitted,



Norman P. Soloway
Attorney for Applicant
Reg. No. 24,315

CERTIFICATE OF EXPRESS MAILING

Express Mail Mailing Label No. EF408018136US Date of Deposit August 16, 2001

I hereby certify that this paper and the papers listed thereon are being deposited with the United States Postal Service "Express Mail Post Office to Addressee" service under 37 CFR 1.10 on the date indicated above and is addressed to the BOX PCT, Assistant Commissioner of Patents, Washington, DC 20231.

Signature of Person Mailing Kimberly Good

Name of Person Mailing Kimberly Good

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MARKED VERSION OF CLAIMS SHOWING CHANGES MADE:

6. (Amended) The hydrogen storage metal alloy according to [any of Claims 1 to 5] Claim 1 wherein the element X is admixed at its atom % concentration, d (at%), ranging within $0 \leq d \text{ (at\%)} \leq 20$, the atomic radius of which is larger than that of Cr but smaller than that of Ti.

8. (Amended) The hydrogen storage metal alloy according to [any of Claims 1 to 7] Claim 1 wherein the element T is admixed at its atom % concentration, e (at%), ranging within $0 \leq e \text{ (at\%)} \leq 10$ and includes at least one or more members selected from the group consisting of Nb, Ta, Mn, Fe, Al, B, C, Co, Cu, Ga, Ge, Ln (a variety of lanthanoid metals), N, Ni, P, and Si.

11. (Amended) The process of Claim 9 [or 10] wherein the predetermined time range at the aforementioned heat treatment is from 1 minutes to 100 hours.

12. (Amended) The process according to [any of Claims 9 to 11] Claim 9 wherein the element ratios are those described in [any of Claims 1 to 8] Claim 1 regarding the hydrogen storage metal alloy products.

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HYDROGEN STORAGE METAL ALLOY AND PRODUCTION THEREOF

5

TECHNICAL FIELD

5 The present invention relates to a hydrogen
storage metal alloy capable of repeatedly carrying
out the absorption and release of hydrogen. In
10 particular, the present invention relates to a BCC
system hydrogen storage metal alloy having
theoretically a high capacity for hydrogen storage.
Further, the present invention especially relates to a
hydrogen storage metal alloy having a highly
15 practicable property, including, for example, not only
quantitatively excellent hydrogen adsorption and
desorption characteristics within practical pressure
ranges and temperature ranges but also a capacity of
adsorbing and desorbing hydrogen in quite great
20 amounts per unit weight, together with a relatively
inexpensive producibility, and to a process for
production thereof.

25

RELATED ART OF THE INVENTION

At present, there have been worried not only
about acid rain due to an increasing NO_x (nitrogen
oxides) but also about the global warming due to an
30 increasing CO₂ in association with an increase in
consumption of fossil fuel such as petroleum. Such
environmental destruction has become a serious problem.
Therefore, our attention has been greatly concentrated
on development and practical application of various
35 kinds of clean energy which is friendly to the earth.
Part of means for developing such a new energy is a
practical application of hydrogen energy. Hydrogen is

5 a constituent element of water inexhaustibly present on the earth and can be not only produced using various kinds of primary energy but also utilized as fluid energy in place of conventionally used petroleum without the risk of destroying the environment because its product by combustion is only water. In addition, unlike electricity, it has excellent characteristics such as its relatively easy storage.

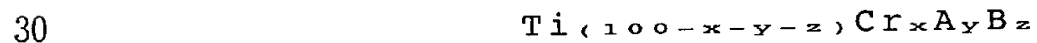
10 In recent years, therefore, investigation has been actively conducted involving hydrogen storage metal alloys as media for storage and transport of hydrogen, and their practical application has been expected. Such hydrogen storage metal alloys are metals/alloys which can absorb and release the
15 hydrogen under an appropriate condition and, by the use of such alloys, it is possible to store the hydrogen not only at lower pressure but also at higher density as compared to the case of the conventional hydrogen cylinders. In addition, the hydrogen volume
20 density thereof is nearly equal to or rather more than that of liquid or solid hydrogen.

These hydrogen storage metal alloys which have been practically used until now are AB₅ alloys such as LaNi₅ and AB₂ alloys such as TiMn₂, but their
25 hydrogen absorbing capacity is still insufficient. In recent years, for example, as proposed in Japanese Unexamined Patent Publication (Kokai) No. 10-110225 (JP, A, 10-110225), metals/alloys (the metal includes V, Nb, Ta, etc. and the alloy does TiCrV alloys, etc.)
30 each having a body-centered cubic (hereinafter, referred to as "BCC") structure have been mostly investigated because the number of hydrogen absorbing sites in the crystal lattice is great in the BCC structure and the hydrogen absorbing capacity is as
35 large as $H/M = \text{ca. } 2$ wherein H is occluded hydrogen and M is a constituent element for the alloy (about 4.0 wt%, i.e., it is huge, in alloys of V, etc. having an

atomic weight of around 50).

With regard to alloys wherein Ti and Cr are comprised, it has been reported as follows: as suggested in JP, A, 10-110225, when for alloys comprised of only Ti and Cr the admixture ratio of the constituent metals is brought to such an extent that it will be conductible to absorb and release hydrogen at a practicable temperature and pressure (i.e., the atomic ratio of Ti is set at $5 < \text{Ti (at\%)} < 60$), as also apparent from FIG. 2 (phase diagram for the Ti-Cr binary alloy), a temperature range for forming a BCC structure becomes very narrow between a melting point of the alloy and a temperature at which a C14 crystal structure is formed. Consequently, other C14 crystal structure phases which are different from BCC are formed at 90 wt% or more in the alloy and it is very difficult to produce the BCC. Therefore, V is admixed as an element highly capable of forming BCC together with both Ti and Cr in order to attain the BCC structure in a more stable fashion and at a lower temperature. As a result, the aforementioned TiCrV alloys have been produced wherein, unless the amount of V is at least 10% or more, it is difficult to form the BCC as their main phase even by application of heat-treatment whereby no good hydrogen adsorption and desorption characteristics are obtainable.

A Ti-Cr based alloy (comprised of 5 or more elements) having the formula:



wherein A is one member selected from V, Nb, Mo, Ta and W, B is two or more members selected from Zr, Mn, Fe, Co, Ni and Cu, and its crystalline structure is BCC is disclosed in Japanese Unexamined Patent Publication (Kokai) No. 7-252560 (JP, A, 7-252560) and it is pointed out therein that the aforementioned admixture

of 5 or more elements is essential for acquiring the
aforementioned BCC.

However, there are still problems: since V to
be admixed with the aforementioned alloy has an atomic
weight approximately similar to that of Ti or Cr, it
may be admixed at an elevated quantity without
reducing its hydrogen storage capacity per unit weight
of the alloy product so much, but because it is very
expensive, especially a highly pure material (99.99 %
purity) employed for such an alloy is extremely
expensive, the price of the alloy product results in a
very high level, whereby alloy costs will increase for
absorbing and storing an equal amount of hydrogen.

Therefore, for inexpensive alloys free of
using precious V, Mo-Ti-Cr and W-Ti-Cr alloys are
proposed wherein Mo or W is admixed as, like V, an
element highly capable of forming BCC with both Ti and
Cr. However, for these Mo and W, as suggested in
Japanese Unexamined Patent Publication (Kokai) No. 10-
121180 (JP, A, 10-121180), it has been reported as
follows: BCC is neither formed in such alloys even by
application of heat-treatments when at 0 at% of Mo
and/or W nor obtainable as the main phase when admixed
at a low level thereof whereby no good hydrogen
absorption and desorption characteristics will appear.
Accordingly, there are also problems: when the amounts
of Mo and W to be admixed increase, the hydrogen
absorbing capacity per unit weight of such alloys will
lower because of their large atomic weight, and in
case where these hydrogen storage metal alloys are
used as energy sources for automobiles, bicycles, etc.
in the form of hydrogen gas storage tanks and nickel
hydrogen batteries, including fuel batteries, when an
attempt is made at attaining a necessary electric
power and hydrogen-supplying capacity, their weights
would increase.

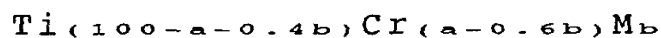
In view of the foregoing points, the present inventors have paid much attention to the aforementioned problems and, as a result, succeeded in the present invention. An object of the present invention is to provide (1) a hydrogen storage metal alloy which is (i) producible in the aforementioned form having BCC main phases even if the level of precious V or Mo and W which each lead to a decrease in hydrogen absorbing capacity per unit weight is made null or as minimal as possible, also (ii) excellent in view of its cost and hydrogen absorbing capacity per unit weight and (iii) highly practicable and (2) a process for producing the same.

SUMMARY OF THE INVENTION

In order to solve the aforementioned problems, the present invention provides a novel hydrogen storage metal alloy for adsorption and desorption of hydrogen. According to the present invention, the novel hydrogen storage metal alloy has the following characteristics:

(1) it has as its main phase a body-centered cubic structure-type phase enabling the adsorption and desorption of hydrogen, and

(2) it has a composition of the following general composition formula:



wherein M is vanadium (V), provided that $20 \leq a \text{ (at\%)} \leq 80$, and $0 \leq b \text{ (at\%)} \leq 10$.

Such characteristics lead to the following: the level of V can be brought to either not more than 10 at% or 0, thereby enabling the amount of necessary precious V to be reduced or nullified, with the result

that hydrogen storage metal alloys thus obtained will become inexpensive, provided that other elements can be optionally admixed as long as their admixture does not have a great influence on the aforementioned properties of the hydrogen storage metal alloys.

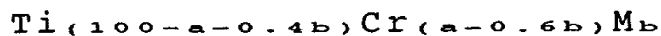
It is preferred that the hydrogen storage metal alloys of the present invention are those wherein the atom % (at%) of element, V, contained in the metal alloy may be within a range of 6 ± 2 at%.

As a result thereof, the alloys can have a higher hydrogen storage capacity per unit weight within a V level range of not more than 10 at% as aforementioned.

According to the present invention, the novel hydrogen storage metal alloy also has the following characteristics:

(1) it has as its main phase a body-centered cubic structure-type phase enabling the adsorption and desorption of hydrogen, and

(2) it has a composition of the following general composition formula:



wherein M is at least a member selected from molybdenum (Mo) and tungsten (W), provided that $20 \leq a$ (at%) ≤ 80 , and $0 \leq b$ (at%) < 5 .

Such characteristics lead to the following:
(1) the level of Mo or W can be brought to either less than 5 at% or 0, thereby enabling the reduction of a hydrogen storage capacity per unit weight, depending on an increase in weight of the resultant alloy, to be minimized or nullified; and (2) since the alloys contain no precious V, they can also be produced inexpensively, provided that other elements can be optionally admixed as long as their admixture does not affect greatly the aforementioned properties of the

hydrogen storage metal alloys.

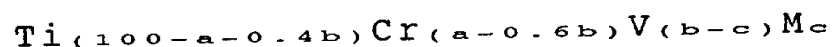
It is preferred that the hydrogen storage metal alloys of the present invention are those wherein the atom % (at%) of Mo and/or W, contained in the metal alloy may be within a range of 3 ± 1.5 at%.

As a result thereof, the alloys can have a higher hydrogen storage capacity per unit weight within a Mo and/or W level range of less than 5 at% as aforementioned.

According to the present invention, the novel hydrogen storage metal alloy also has the following characteristics:

(1) it has as its main phase a body-centered cubic structure-type phase enabling the adsorption and desorption of hydrogen, and

(2) it has a composition of the following general composition formula:



wherein M is at least a member selected from molybdenum (Mo) and tungsten (W), provided that $20 \leq a$ (at%) ≤ 80 , $0 < b$ (at%) < 10 , and $0 < c$ (at%) < 5 .

Such characteristics lead to the following: part of the content of precious V can be replaced with at least one member selected from Mo and W which are each potentially capable of forming a BCC structure together with Ti and Cr in the same manner as V, thereby enabling not only the cost to be relatively low but also a decrease in hydrogen storage capacity per unit weight, brought about by the incorporation of Mo or W, to be limited to a relatively minor one, with the result that the hydrogen storage metal alloys can be produced which come to an excellent balance between the cost and the hydrogen storage capacity per unit weight and become advantageously practicable, provided that other elements can be optionally admixed as long

as their admixture does not affect greatly the
aforementioned properties of the hydrogen storage
metal alloys.

It is preferred that the hydrogen storage
5 metal alloys of the present invention are those
wherein an element, X, having an atomic radius greater
than that of Cr but smaller than that of Ti may be
contained at its atom % concentration, d (at%),
ranging within $0 \leq d \text{ (at\%)} \leq 20$.

10 As a result thereof, the element X can be
admixed the atomic radius of which is larger than that
of Cr but smaller than that of Ti, thereby inhibiting
the formation of a C14 (Laves phase) structure so as
to extend a temperature range for forming a BCC
15 structure phase in place of the aforementioned C14
(Laves phase) structure, with the result that the
hydrogen storage metal alloys can be produced with the
BCC structure phase in a stable fashion even at low
levels of V, Mo and W, which each have a potent BCC
20 structure-forming capability with both Ti and Cr.

It is preferred that the hydrogen storage
metal alloys of the present invention are those
wherein the element, X, may include at least one or
more members selected from the group consisting of Al,
25 Ge, Ga, Si, Au and Pt.

As a result thereof, the selected elements
have an excellent capability of forming a metal alloy
with Ti and Cr and therefore preferable to be employed
for the aforementioned element X.

30 It is preferred that the hydrogen storage
metal alloys of the present invention are those
wherein an element, T, may be contained at its atom %
concentration, e (at%), ranging within $0 \leq e \text{ (at\%)} \leq 10$,
wherein the said element T includes at least one or
35 more members selected from the group consisting of Nb,
Ta, Mn, Fe, Al, B, C, Co, Cu, Ga, Ge, Ln (a variety of
lanthanoid metals), N, Ni, P, and Si.

As a result thereof, the element T can be admixed, thereby enabling a plateau pressure at which the resultant hydrogen storage metal alloys can absorb and release hydrogen to be appropriately controlled.

5 According to the present invention, a process for producing the hydrogen storage metal alloy has the following characteristics:

for the production of hydrogen storage metal alloys having as the main phase a body-centered cubic structure-type phase enabling the adsorption and desorption of hydrogen, and comprising the steps of:

(1) melting a starting alloy brought to a predetermined element ratio to form a uniform heat (melting step),

15 (2) keeping the homogenized alloy heat at a temperature within a range just below the melting point of the alloy for a predetermined time (heat treatment), and

(3) rapidly cooling the alloy after the heat treatment (quenching step).

20 Such characteristics enable the production of hydrogen storage metal alloys having as the main phase a BCC-type phase with regard to not only a Ti-Cr binary alloy, which is presumed to be hardly produced, but also an alloy wherein V, Mo and W are contained at low levels.

It is preferred that melting and solidification may be carried out repeatedly predetermined times at the aforementioned melting step in the hydrogen storage metal alloy-producing process according to the present invention.

30 As a result thereof, such repeated melting and solidifying treatments enable us not only to produce alloys having an improved uniformity wherein a BCC-type structure phase is formed at a higher rate but also to prevent the occurrence of a spinodal decomposing composition as much as possible.

It is preferred that the predetermined time range at the aforementioned heat treatment is from 1 minute to 100 hours in the hydrogen storage metal alloy-producing process according to the present invention.

When a time range for the heat treatment step is 1 minute or less, it is impossible to form sufficiently a BCC-type structure and when 100 hours or more, treating costs increase due to heating for a long time. As a result thereof, the inventive time range leads to the excellent formation of the BCC structure with minimizing an increase in the treating cost.

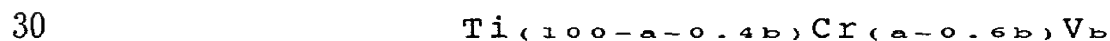
It is preferred that the element ratios are those described in any of Claims 1 to 8 regarding the aforementioned hydrogen storage metal alloy products obtained by the hydrogen storage metal alloy-producing process according to the present invention.

As a result thereof, alloys wherein the main phase is a BCC-type structure can be produced in a stable fashion from each alloy having a highly practicable composition according to any of Claims 1 to 8.

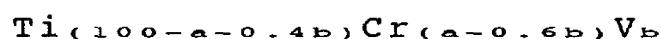
Described below are bases of the selected compositions for hydrogen storage metal alloys according to the present invention. FIG. 2 depicts a Ti-Cr binary system phase diagram in connection with the present invention. As seen in FIG. 2, the BCC phase is present throughout all composition ranges of Ti and Cr over 1643K (1370°C). In light of the atomic radius of Ti (0.147 nm) greater than that of Cr (0.130 nm), when the level of Ti increases and the level of Cr lessens, the alloy will increase its BCC phase lattice constant but lower its plateau pressure. The plateau pressure of the hydrogen storage metal alloy may vary depending on its alloy-working temperature. It is preferred that the Ti/Cr ratio may vary in order to

acquire a desired working temperature and consequently a suitable Ti/Cr ratio can be selected. Although a starting composition is brought to the extent of $Ti_{40}Cr_{60}$ in order to acquire a suitable plateau pressure at $40^{\circ}C$ (313K) in examples as described hereinbelow, the present invention is not limited to. The plateau pressure of such hydrogen storage metal alloys not only varies depending on their alloy-working temperature but also is controllable by varying a Ti/Cr ratio for Ti-Cr-M hydrogen storage metal alloys including Ti-Cr alloys and Ti-Cr-V alloys. The plateau pressure remarkably rises when the Cr level "a" exceeds 80 at% while the plateau pressure becomes extremely low when it is below 20 at%, thereby leading to a poor practicability. Accordingly, it is preferred that the Ti/Cr ratio may be selected which is suited for a desired working temperature within a range of $20 \leq a \text{ (at\%)} \leq 80$.

Although the addition of V to such Ti-Cr binary alloys is effective in facilitating the formation of their BCC type structure as aforementioned, an excessive admixture of V leads to a decrease in their hydrogen adsorption and desorption characteristics as shown in FIG. 5, and when the V level is over about 10 at%, the admixture of such precious V will not make any sense. In light of the foregoing points, it is derived that the fundamental formula:

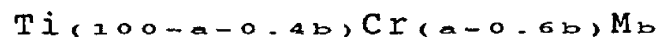


will be within a range of $0 \leq b \text{ (at\%)} \leq 10$. Further, the addition of substituent element T to such alloys having the fundamental formula:



is effective in adjusting the plateau pressure wherein T is at least one or more elements selected from the group consisting of Nb, Ta, Mn, Fe, Al, B, C, Co, Cu, Ga, Ge, Ln (a variety of lanthanoid metals), N, Ni, P, and Si, and an amount of substituents is $0 \leq c \text{ (at\%)} \leq 10$.

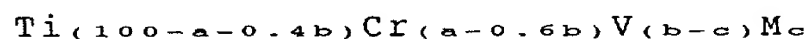
Although Mo and W elements each have a great BCC structure-forming property to Ti-Cr binary alloys and the admixture of Mo or W with the Ti-Cr binary alloy is effective in facilitating the formation of the BCC structure, an excessive admixture of Mo and/or W will lead to an increase in density for hydrogen storage metal alloy products because such Mo and W are heavy elements with a relatively large atomic weight. As seen in FIGS. 9 and 10, when their level exceeds about 5 at%, the hydrogen adsorption and desorption characteristics which reach to the maximum will be led to a significant decrease. In light of the foregoing points, the fundamental formula:



wherein a is $20 \leq a \text{ (at\%)} \leq 80$, b is $0 \leq b \text{ (at\%)} < 5$ and M is at least one element selected from Mo and W, is derived. The admixture of substituent element T with the resultant metal alloys is also effective in adjusting the plateau pressure in the same fashion as aforementioned, provided that T is at least one or more elements selected from the group consisting of Nb, Ta, Mn, Fe, Al, B, C, Co, Cu, Ga, Ge, Ln (various lanthanoid metals), N, Ni, P and Si, and an amount of substituents is $0 \leq c \text{ (at\%)} \leq 10$.

Further, although element V has an atomic weight approximately equivalent to that of Ti or Cr and is precious, even a large quantity of its substituent leads to a less increase in molecular weight for alloy products whereby there is an

advantage that an amount of occluded hydrogen per unit weight will not be reduced much. In contrast, since Mo and W each have a great BCC structure-forming property to Ti-Cr binary alloys, the admixture of Mo and/or W with the Ti-Cr binary alloy is effective in facilitating the formation of BCC in alloy products. However, an excessive admixture of Mo and W will lead to a decrease in hydrogen adsorption and storage characteristics because of heavy elements each having a large atomic weight. Hence, to make better use of both the advantages, a novel composition is invented wherein part of precious V is replaced with Mo and/or W, i.e., an alloy composition of the following fundamental formula:



wherein $20 \leq a \text{ (at\%)} \leq 80$, $0 \leq b \text{ (at\%)} \leq 10$, $0 \leq c \text{ (at\%)} < 5$, and M is at least one element selected from Mo and W, is greatly practicable in view of its cost and its occluded hydrogen quantity as well as its BCC structure-forming capability. As before, the admixture of substituent element T with such a composition is also effective in adjusting the plateau pressure wherein T is at least one or more elements selected from the group consisting of Nb, Ta, Mn, Fe, Al, B, C, Co, Cu, Ga, Ge, Ln (various lanthanoid metals), N, Ni, P and Si.

Alloys having a low level of these elements and Mo or W are hardly formed in the structure of BCC as pointed out in the prior art. As apparent from the phase diagram of a Ti-Cr binary alloy (FIG. 2), this is attributable to the fact that a temperature range for affording the BCC structure is too narrow throughout the Ti-Cr admixture ratios wherein temperature and pressure ranges at which the hydrogen storage metal alloy can work will be within

practicable values, i.e., at the Cr level of 20 to 80 at%.

As seen in the aforementioned phase diagram (FIG. 2), however, when the level of Cr is gradually reduced from 60 at% (it has the same meaning as the level of Ti gradually increase from 40 at%), a temperature range eligible for giving a BCC structure would expand. This is presumably attributed to the following: since the Laves phase is represented by a composition of a AB_2 type and the atomic radius ratio of A to B ($r_A : r_B$) = about 1.225 : 1 is necessary for forming an ideal geometric structure in such a composition while the atomic radius ratio of Ti to Cr, both of which are used according to the present invention, is 1.13 : 1, which is far different from the above ideal value and unsuitable for forming the ideal Laves phase structure, Ti will quantitatively increase, and invade B sites in apparently more quantities whereby consequently the atomic radius ratio at A sites will become close to that at B sites, thereby inhibiting the formation of Laves phases.

Now, by developing such ideas, when an element having an atomic radius smaller than that of the A site but larger than that of the B site is admixed therewith for substitution, not only the penetration of the substituent element into the A site can inhibit the formation of Laves phase but also that into B site can it similarly.

Hence, it has been thought that there is a possibility of enabling a BCC formation in alloy products similarly to the above V case as well as the Mo or W case and therefore element X (its atomic radius is smaller than that at the A site (Ti) but larger than that at the B site (Cr)) can be admixed with the alloy to expand a temperature range eligible for forming BCC whereby a hydrogen storage metal alloy may be produced with a BCC structure in a more stable

fashion.

The element X, the atomic radius of which is smaller than that at the A site (Ti) but larger than that at the B site (Cr), includes, for example, at least one or more elements selected from the group consisting of Al (0.143 nm), Si (0.132 nm), Ga (0.141 nm), Ge (0.137 nm), Au (0.146 nm) and Pt (0.139 nm) in view of Ti atomic radius with 0.147 nm and Cr with 0.130 nm.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a flow chart illustrating a process for producing the hydrogen storage metal alloy according to an embodiment of the present invention.

FIG. 2 depicts a Ti-Cr binary system phase diagram.

FIG. 3 is an X-ray diffraction pattern of as heat-treated (at 1400 °C for 1 hour) alloy



FIG. 4 is a graph showing hydrogen absorption and desorption characteristics (at 40 °C) for as heat-treated (at 1400°C for 1 hour) alloy



FIG. 5 is a graph showing the relationship of admixed amounts of V versus hydrogen absorption and desorption characteristics for a Ti-Cr-V alloy.

FIG. 6 is an X-ray diffraction pattern of as heat-treated (at 1400 °C for 1 hour) alloy

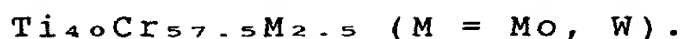


FIG. 7 is a graph showing hydrogen absorption and desorption characteristics (at 40 °C) for as heat-treated (at 1400°C for 1 hour) alloy $Ti_{40}Cr_{57.5}Mo_{2.5}$.

5 FIG. 8 is a graph showing hydrogen absorption and desorption characteristics (at 40 °C) for as heat-treated (at 1400°C for 1 hour) alloy $Ti_{40}Cr_{57.5}W_{2.5}$.

10 FIG. 9 is a graph showing the relationship of admixed amounts of Mo versus hydrogen absorption and desorption characteristics for a Ti-Cr-Mo alloy.

15 FIG. 10 is a graph showing the relationship of admixed amounts of W versus hydrogen absorption and desorption characteristics for a Ti-Cr-W alloy.

20 FIG. 11 is an X-ray diffraction pattern each of as heat-treated (at 1400 °C for 1 hour) alloys $Ti_{37.5}Cr_{60}V_{2.5}$ and $Ti_{37.5}Cr_{60}Mo_{1.25}V_{1.25}$.

FIG. 12 is an X-ray diffraction pattern each of as prepared by melting and as heat-treated alloys $Ti_{40}Cr_{60}$.

25 FIG. 13 is an X-ray diffraction pattern each of as heat-treated (at 1400 °C for 1 hour) alloy $Ti_{42.5}Cr_{57.5}$ and as heat-treated (at 1400 °C for 2 hours) alloy $Ti_{40}Cr_{60}$.

30 FIG. 14 is a graph showing hydrogen absorption and desorption characteristics (at 40 °C) for as heat-treated alloy $Ti_{42.5}Cr_{57.5}$.

35 FIG. 15 is an X-ray diffraction pattern of as heat-treated (at 1400 °C for 1 hour) alloy $Ti_{40}Cr_{57.5}Al_{2.5}$.

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FIG. 16 is a graph of hydrogen absorption and desorption characteristics (release curve, 40 °C, 5th cycle) upon application of differential temperature method to alloy

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$V_xTi_{37.5}Cr_{62.5-x}$.

BEST MODE FOR CARRYING OUT THE INVENTION

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Described below are the hydrogen storage metal alloys of the present invention and processes of the production of the said metal alloys in detail, relying on experiments conducted by the present inventors.

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First, FIG. 1 is a flow chart showing a preferred embodiment of the process for producing the hydrogen storage metal alloys according to the present invention. Such a process has been applied to the production of hydrogen storage metal alloys used in the experiments conducted by the present inventors as described hereinbelow.

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In this process of the production of hydrogen storage metal alloys, each constituent element for a desired hydrogen storage metal alloy (for example, each of Ti, Cr and V in case of producing $Ti_{37.5}Cr_{60}V_{2.5}$) was weighed at an amount corresponding to each composition ratio so as to bring the total weight of a resultant ingot to 12.5g.

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Each individual metal thus weighed was thrown into an arc melting plant (not depicted), subjected to repeated treatments (melting-stirring $\leftarrow \rightarrow$ solidification) predetermined times (which may vary depending on the number of constituent elements in experiments but be usually approximately 4 to 5 times) in an argon atmosphere of about 40 kPa with scrupulous care to elevate a uniformity and the resultant homogenized ingot was then maintained at a temperature

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35

region just lower than the melting point of its melt for a predetermined time to accomplish the heat treatment.

Since a temperature region at which BCC forms are produced is present at an area just below the melting temperature owned by an alloy having a target composition as shown in the above FIG. 2 (phase diagram), the heat-treatment may be preferably effected at such a temperature region at which the BCC is produced and just below the melting temperature. For example, in case of a composition containing about 60 at% of the aforementioned element Cr, although the heat-treatment is preferably effected by holding the molten alloy at about 1400 °C, it is preferable to select a suitable heat-treating temperature from temperature areas at which a target alloy is produced in the form of BCC and just below the melting temperature of the target alloy, depending on its alloy composition. Among temperature areas at which the BCC is produced and just below the melting temperature thereof, it should be noted that it will take a longer time to accomplish the heat-treatment when the treatment temperature is too low (not more than about 1000 °C) while it will take only a short time but heating will cost us too much when it is too high. Therefore, by taking the foregoing points into account, it is preferable to select a heat-treating temperature.

When a heat-treating time is too short, it will be impossible to accomplish the formation of sufficient BCC structure phases while when it is too long treating, costs will rise dependently on heating for a long period. Although it is therefore preferred that the heat treatment can be conducted for a period suitably selected on the basis of a selected heat-treating temperature, it may be carried out preferably for a period of from 1 min to 100 hours, more

preferably from 10 min to 24 hours, in the embodiment, from 1 to 2 hours.

In the Examples, alloys per se were subjected to the aforementioned heat treatment after melting ingots without making any shapes. Since such a process does not require that cooled alloys are re-heated but enables us to produce efficiently alloy products having a BCC structure phase, it is preferable but the present invention is not limited to. For example, it may be preferred that molten alloys are shaped once by methods such as strip casting, mono roll casting and atomizing methods to afford plates, ribbons or powders, then cooled and the resultant alloys each having either the BCC phase + the Laves phase or the Laves phase alone are subjected to the aforementioned heat treatment so as to form the BCC structure phase as the main phase.

Among these alloys, alloys (ingots) heat treated to an extent that the BCC structure phase takes place as the main phase are rapidly cooled by dipping into ice water to give alloy products in the form of holding the above BCC structure phase.

Although the aforementioned rapid cooling has been carried out by dipping into ice water, the present invention is not limited to and any can be arbitrarily selected for these cooling methods. However, since the volume ratio of BCC structural phase varies depending on cooling rates and a slow cooling rate leads to a decrease in BCC structural phase volume ratio, it is desired that the alloy is rapidly cooled preferably at a cooling rate of more than 100K/sec.

Although the alloys of the present invention have a composition apt to induce a spinodal decomposition readily, it is defined that they are acceptable to the extent there is an unavoidable formation because spinodal decomposing tissues cause

deterioration of alloy's hydrogen absorption and desorption characteristics.

Hereinbelow not only it has been examined and ascertained whether or not the BCC structural phase is produced as the main phase by the aforementioned production processes for each composition but also experimental results are shown which support bases of the above selected compositions.

An X-ray diffraction pattern of an alloy product obtained by heat-treating $V_xTi_{37.5}Cr_{62.5-x}$ alloy at 1400 °C for 1 hour is shown in FIG. 3. As apparent from FIG. 3, even when V which is presumed to be hardly utilizable in the prior art as aforementioned is admixed at 2.5 at%, the BCC takes place as the main phase and when V is set to 5 at% and 7.5 at%, respectively, the alloys are produced in the form of a BCC mono phase.

The fact that each alloy as shown in FIG. 3 has the BCC structure reflects on its hydrogen absorption and desorption characteristics as shown in FIG. 4. Thus, it has been found that the BCC mono phase alloys wherein V is contained at 5 at% and 7.5 at% can absorb and release hydrogen at about 2.8 wt% which is approximately equal to or more than the amount achieved by the prior art alloy containing 10 at% or more of V. Further, it has been found that even the alloy wherein V is contained at 2.5 at% can absorb and release hydrogen at about 2.6 wt % which is approximately equal to the amount achieved by the prior art alloy containing 10 at% or more of V.

This is that the admixture of V with the Ti-Cr binary alloy derives an increase in BCC structural phase volume ratio, whereby an amount of occluded hydrogen increases over that attained by the Ti-Cr binary alloy. Thus, it has been found that V is an element greatly apt to produce a BCC form and effective for bringing advantageous hydrogen

absorption and desorption characteristics owned by the BCC phase into the Ti-Cr binary alloys. The hydrogen absorption and desorption characteristics affected by amounts of admixed V in connection with these Ti-Cr-V alloys were examined. The results are shown in FIG. 5.

The results as shown in FIG. 5 are unexpected ones. When an amount of the admixed element V is brought to 10 at% or more, which was considered to be preferable in the prior art, it is ascertained that target alloys to be produced are improved for their capability of forming the BCC phase and consequently alloy products having the BCC phase become manufacturable in a stable fashion according to an increase in amount of element V to be admixed; nevertheless, it results in their hydrogen storage capacity per unit weight equal to or less than that of V-free Ti-Cr binary alloys (without other materials). It is therefore apparent that their hydrogen storage capacity per unit weight reaches to a maximum at a V admixture amount of not more than 10 at%, especially 6 ± 2 at%, contrary to conventional understanding in the prior art. Accordingly, it is found that an amount of admixed V can be set to such a region so as to not only prevent an increase in cost for alloys due to production by unnecessarily admixing an excessive amount of precious V but also increase a hydrogen storage capacity per unit weight.

Next, the aforementioned production process has been applied to Ti-Cr-Mo (W) system hydrogen storage metal alloys which have been associated with the aforementioned problems, for example, when Mo and W which are each a heavy element having a high capability of forming BCC against Ti-Cr alloys but a large atomic weight, are admixed in large quantities, the alloy products hardly exert sufficient properties, etc. Each level of Mo and W therein has been examined. The results are described hereinbelow.

An X-ray diffraction pattern each of as heat-treated alloys $Ti_{40}Cr_{57.5}Mo_{2.5}$ and $Ti_{40}Cr_{57.5}W_{2.5}$ is shown in FIG. 6. It is found from the X-ray diffraction pattern as shown in FIG. 6 that, although Mo is admixed at a small amount, i.e., at 2.5 at% only, the resultant alloy products are in the form of a BCC mono phase. For W, BCC phases are produced as the main phase though a few Laves phases are present.

Further, hydrogen adsorption and desorption characteristics of as heat-treated alloy $Ti_{40}Cr_{57.5}Mo_{2.5}$ are shown in FIG. 7. It has been noted therefrom that an amount of hydrogen occluded thereby is derived to an extent of about 2.9 wt% close to a value of 3 wt% corresponding to the maximal capacity which is considered to be intrinsically owned by the Ti-Cr binary BCC phase.

From these results, it has been found that Mo can be admixed even at a smaller amount than V so as to produce almost the BCC mono phase. Thus, it is noted that, as compared to the foregoing Ti-Cr-V alloys, an amount of additives can be reduced whereby a good property has been achieved.

Hydrogen adsorption and desorption characteristics of as heat-treated alloy $Ti_{40}Cr_{57.5}W_{2.5}$ are also shown in FIG. 8. Similarly to the aforementioned Mo, the W-substituted alloys come to be in the form of a BCC mono phase and their hydrogen adsorption capacity reaches to about 2.7 wt % or more. When W is admixed at an amount equal to that for Mo or V, the resultant alloy products come to decrease their maximal hydrogen adsorption capacity slightly because W has a larger atomic weight.

The hydrogen adsorption and desorption characteristics which an amount each of additives, Mo and W, affects for such heat-treated Ti-Cr-Mo and Ti-Cr-W alloys are shown in FIGs. 9 and 10, respectively. When the additive element is Mo, it has been found

that the admixture of Mo at a small amount leads to an increase in hydrogen adsorption capacity, i.e., the hydrogen adsorption capacity becomes maximal at about 3 ± 1.5 at% of the additive Mo while the hydrogen adsorption capacity comes to be gradually decreased in the 5 at% or more Mo regions which have been presumed to be preferable in the prior art. It has also been found that when Mo is admixed at 10 at% or more, the hydrogen adsorption capacity comes to lower less than that of heat-treated Mo-free Ti-Cr alloys. When the additive element is W, it has been observed that the resultant alloy products have a tendency similar to that for the aforementioned Mo. It has thus been found that the admixture of W at a small amount leads to an increase in hydrogen adsorption capacity, i.e., the hydrogen adsorption capacity becomes maximal at about 3 ± 1.5 at% of the additive W while the hydrogen adsorption capacity comes to be gradually decreased in the 5 at% or more W regions which have been presumed to be preferable in the prior art. It has also been found that when W is admixed at 6 at% or more, the hydrogen adsorption capacity comes to lower less than that of heat-treated W-free Ti-Cr alloys.

Hence, these elements Mo and W are preferably admixed therewith at a micro amount so as to increase occupied volume ratios of the BCC phase occurred in such Ti-Cr binary alloys. It is found that Mo and W, as compared for their capability of forming BCC in Ti-Cr alloys, have each a tendency to enable the BCC-occupied volume ratio to increase even upon their admixture at a smaller amount than V's. It is also found that amounts per unit weight of occluded hydrogen therein come to decrease when they are admixed at an excessive amount.

Although each of Mo and W is admixed alone in order to clarify the efficacy of individual admixed elements in the foregoing embodiments, the present

invention is not limited to. It is preferred that one of two elements Mo and W may be admixed therewith in combination with the other. For amounts of the admixed elements in this instance, it is preferable that a
5 total amount of admixed elements Mo and W may be less than 5 at%.

As aforementioned, V has an atomic weight approximately equivalent to that of Ti or Cr. Although V is precious, molecular weights of alloy
10 products are changed (increased) little even when an amount of substituents increases. Therefore, there are advantages that amounts of occluded hydrogen do not reduce very much. Accordingly, in order to produce BCC mono phase alloys with a high capacity by melting a
15 large amount of alloys followed by rapidly cooling and, if necessary, heat-treatments, it is forecasted that V may be effectively admixed therein in combination with at least one member selected from the aforementioned Mo, W, etc. Thus, for the aforementioned Ti-Cr-V
20 alloys wherein a low level of V is contained, which have been conventionally presumed to be hardly produced in a BCC phase form, efficacies are examined and proved in case where a replacement with Mo partially takes place.

25 An X-ray diffraction pattern each of as heat-treated $Ti_{37.5}Cr_{60}V_{2.5}$ and $Ti_{37.5}Cr_{60}Mo_{1.25}V_{1.25}$ alloys is shown in FIG. 11. Reflections by the Laves phase are observed for the heat-treated alloy $Ti_{37.5}Cr_{60}V_{2.5}$ as shown in FIG. 11 (identical with the pattern for
30 $X=2.5$ as shown in FIG. 3) and the hydrogen adsorption and desorption characteristics only reach to an extent of 2.6 %. However, it has been found that the heat-treated alloy $Ti_{37.5}Cr_{60}Mo_{1.25}V_{1.25}$ wherein part of V elements are replaced with Mo are almost in the form
35 of a BCC mono phase and its hydrogen adsorption and desorption characteristics are improved to be an extent of about 2.7 wt%. In this way, V can be admixed

therewith in combination with Mo(also W) so as to reduce an amount of precious V to be admixed together with a reduction in amounts of Mo (and/or W) to be admixed, with the result that the occupied volume ratio of BCC phases will increase together with these admixtures, thereby leading to an increase in hydrogen adsorption capacity. Therefore, it can be said that the admixture of V in combination with Mo (and/or W) is a preferable technique for producing inexpensive hydrogen storage metal alloys with a high capability of absorbing and storing hydrogen.

It has been proved that, as aforementioned, by using the above production process characterized in the heat-treatment of the present invention the BCC phase is produced as the main phase at an area (wherein an amount each of additives V, Mo, W, etc. is extremely slight) approximately close to the Ti-Cr binary alloy which is conventionally presumed to produce no BCC phase in the prior art and the resultant alloys exert excellent hydrogen adsorption and desorption characteristics. Accordingly, it has been found that there is a possibility of acquiring excellent hydrogen adsorption and desorption characteristics with regard to Ti-Cr binary alloys, i.e., alloys free of additives such as V, Mo and W but formed from only Ti-Cr, wherein it has been presumed in the prior art that the BCC would be hardly formed as the main phase and accordingly no good hydrogen absorption and release would be achieved. Next, occurred phases and hydrogen adsorption and desorption characteristics are examined for such Ti-Cr binary alloys.

An X-ray diffraction pattern each of as prepared by melting (as cast) and as heat-treated at 1673K (kept at 1400 °C for 1 hr followed by water-cooling) $Ti_{40}Cr_{60}$ is shown in FIG. 12.

It is apparent from the X-ray diffraction

patterns as shown in FIG. 12 that the BCC is produced as the main phase. Next, in order to have a try at improving on Ti-Cr binary alloys for forming the BCC mono phase, researches are conducted on alloy compositions and conditions for heat-treatments. An X-ray diffraction pattern each of $Ti_{42.5}Cr_{57.5}$ alloy materials heat-treated at 1673K (maintained at 1400 °C for 1 hr followed by cooling with water) and $Ti_{41}Cr_{59}$ alloy materials heat-treated at 1673K wherein a time range for heat-treatment was 2 hours (maintained at 1400 °C for 2 hr followed by cooling with water) is shown in FIG. 13. It is apparent from this drawing that both alloys have the main phase of the BCC. Especially the former is produced in the form of a BCC mono phase regardless of the same heat-treating conditions as for the alloy as shown in FIG. 12. These results indicate that the present invention solves the problems (it has been reported in JP, A, 10-121180, Japanese Unexamined Patent Publication (Kokai) No. 10-158755 (JP, A, 10-158755), Japanese Unexamined Patent Publication (Kokai) No. 11-106859 (JP, A, 11-106859) that it was difficult to produce Ti-Cr binary alloys in the form of a BCC mono phase) by adjusting the alloy compositions to most suitable ones, fitting treating time, etc.

From these experimental Ti-Cr alloy results (see: FIGs. 12 and 13), it is suggested that the Laves phase formation is inhibited easier in alloy $Ti_{42.5}Cr_{57.5}$ than in alloy $Ti_{40}Cr_{60}$, that is, it is prevented easier in alloys replaced with Ti having a larger atomic radius (0.147 nm) than Cr (0.130 nm). The Laves phase is represented by a composition AB_2 wherein the atomic radius ratio between both atoms A and B, i.e., ($r_A : r_B$) is about 1.225 : 1 in order to have a geometrically ideal structure and also characterized by the composition ratio, A : B, having a range. However, the ratio of Ti atomic radius : Cr

atomic radius is 1.13 : 1, which is unsuitable for forming an ideal Laves phase structure from an initial stage, and the Ti : Cr atom % ratio, which is a target to be researched according to the present invention, is about 1 : 1.5, i.e., a large amount of Ti apparently occupies a B site, whereupon the atomic radius ratio between the A site and the B site also reduces, etc. These are thought to be sources for inducing results different from those reported in the prior art.

Relying on an expansion of these ideas, the following will be expected: when a replacement with an element having an atomic radius smaller than the A site but greater than the B site takes place, even the intrusion of the substituent elements into A sites will lead to the inhibition of the Laves phase formation and that into B sites will as well. That is, it is forecasted that there are elements facilitating the formation of BCC. It is expected that a substitution with an element (including, for example, Al (0.143 nm), Ga (0.141 nm), Ge (0.137 nm) and Pt (0.139 nm), etc.) having an atomic radius greater than Cr (0.130 nm) but smaller than Ti (0.147 nm) will facilitate the formation of the BCC phase.

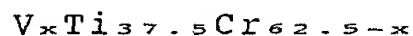
Thus, there has been no report that the Ti-Cr binary alloy was subjected to the formation of a BCC mono phase or the facilitation of a BCC phase formation. This is one of bases supporting the novelty of the present invention. The hydrogen absorption and desorption characteristics of as heat-treated alloy $Ti_{42.5}Cr_{57.5}$ are shown in FIG. 14. Its hydrogen storage capacity is 2.6 wt % or more. Distinctively from Ti-Cr Laves alloys and the like as reported in the prior art, these results evidence that the BCC phase occurring in the Ti-Cr binary alloy has advantageous hydrogen adsorption and desorption characteristics.

In connection with the fact that BCC phases occur in ternary system alloys such as Ti-Cr-V and Ti-Cr-M (M=Mo or W) alloys, which have been disclosed in JP, A, 10-121180, JP, A, 10-158755 and JP, A, 11-106859, the following has been experimentally proved: Ti-Cr-V alloys and Ti-Cr-Mo (W) or Ti-Cr-(V or Mo) alloys according to the present invention are produced in the form of a BCC mono phase or in a BCC main phase form at an extremely micro amount of V, Mo, W, etc. to be admixed (i.e., at a range very close to the Ti-Cr binary alloy), thereby exerting excellent hydrogen adsorption and desorption characteristics. This is attributed to the fact that the BCC phase of such Ti-Cr binary alloys exerts its excellent hydrogen adsorption and desorption characteristics.

An X-ray diffraction pattern each of as heat-treated alloys $Ti_{40}Cr_{60}$ and $Ti_{40}Cr_{37.5}Al_{2.5}$ is shown in FIG. 15. It is apparent that a BCC mono phase is almost formed by replacing part of Cr with Al. This alloy is realized by using $Ti_{42.5}Cr_{57.5}$ alloy rather than $Ti_{40}Cr_{60}$ alloy as shown for Ti-Cr alloys, i.e., by developing the concept that Cr is replaced with Ti having a larger atomic radius than Cr to bring the atomic radius ratio of A to B ($r_A : r_B$) to such an extent that it will facilitate the inhibition of a Laves phase formation and applying Al (0.143 nm) which has an atomic radius larger than Cr (0.130 nm) but smaller than Ti (0.147 nm) and is capable of not only inhibiting the formation of a Laves phase but also facilitating the formation of BCC even irrespective of which of A and B sites is replaced. Elements to be admixed therewith are those which have an action similar to Al and may include Ga, Ge, Si, Pt, Au, etc.

The reason why alloys each having such an alloy composition (such ingredients) can be readily designed is that they are based on Ti-Cr binary BCC alloys in accordance with the present invention,

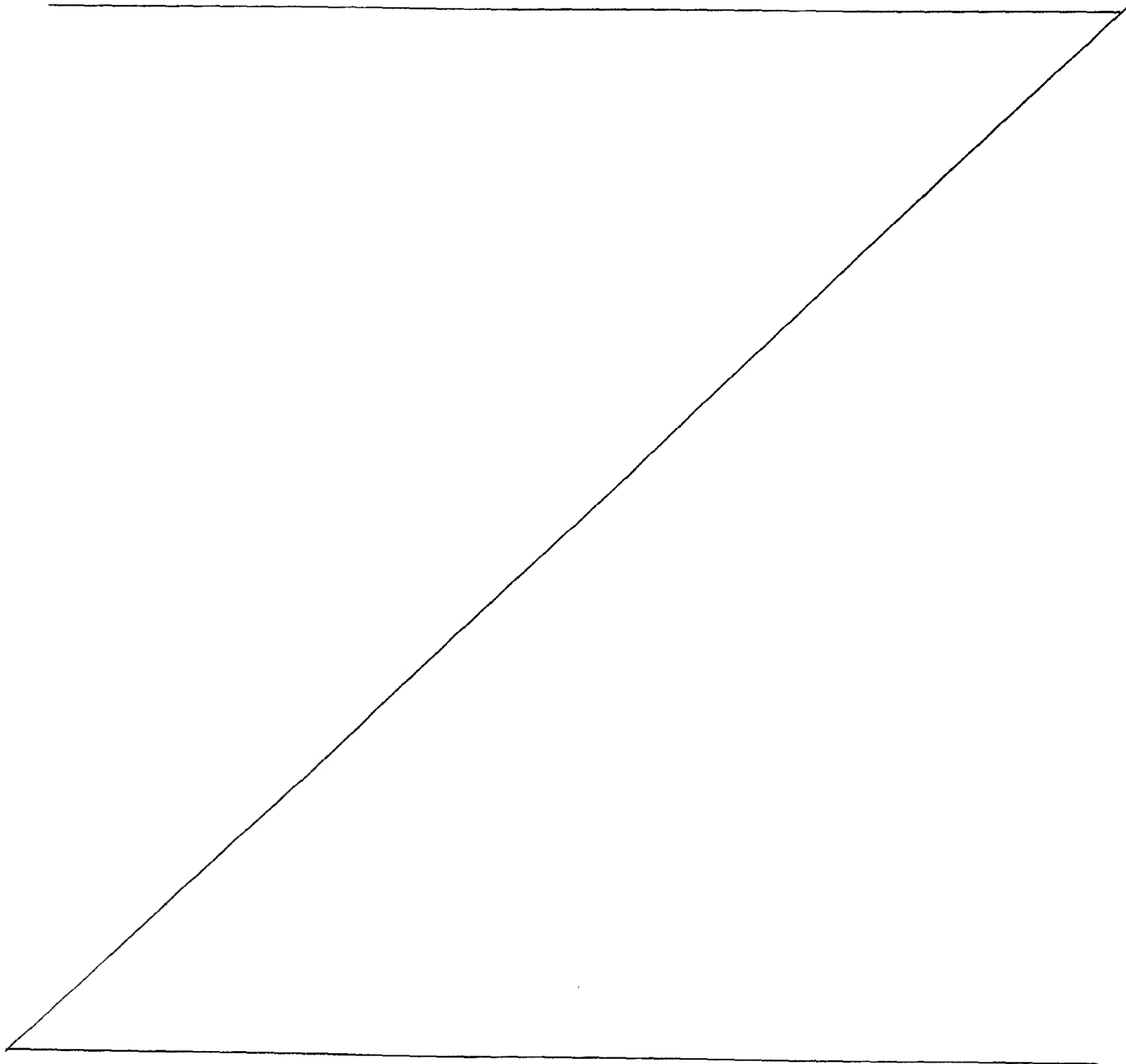
unlike the prior art. It is reported in Japanese Patent Application No. 11-86866 (or 86866/1999) that hydrogen can efficiently be utilized via applications of a difference in temperature, characterized in that
5 body-centered cubic structure hydrogen storage metal alloys each having a two-stage plateau or inclined plateau are subjected to an occlusion of hydrogen at a low temperature followed by an elevation of an alloy working temperature for at least a period of hydrogen
10 release process. In case where the differential temperature method is applied to the aforementioned alloy



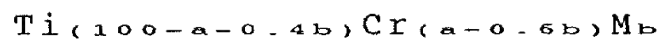
15 its hydrogen absorption and desorption characteristics are shown in FIG. 16. It is apparent that the application of the differential temperature method to the alloys of the present invention will lead to a hydrogen storage capacity of about 3.0 wt %. As
20 compared to FIG. 4, it is observed that the differential temperature method derives an increase in hydrogen storage capacity at about 0.2 wt %, and it is therefore experimentally proved that the differential temperature method is effective for alloys attained by
25 the present invention. Its practicability can also be understood.

What is claimed is:

1. A hydrogen storage metal alloy which has
 - (1) as its main phase a body-centered cubic
5 structure-type phase enabling the adsorption and
desorption of hydrogen, and
 - (2) a composition of the following general
composition formula:



09013625 081604
09013625 081604



wherein M is vanadium (V), provided that $20 \leq a$ (at%) ≤ 80 , and $0 \leq b$ (at%) ≤ 10 .

5

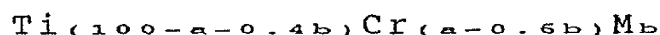
2. The hydrogen storage metal alloy according to Claim 1 wherein a level of the constituent element V contained in the alloy is within a range of 6 ± 2 at %.

10

3. A hydrogen storage metal alloy which has
(1) as its main phase a body-centered cubic structure-type phase enabling the adsorption and desorption of hydrogen, and

15

(2) a composition of the following general composition formula:



20

wherein M is at least a member selected from molybdenum (Mo) and tungsten (W), provided that $20 \leq a$ (at%) ≤ 80 , and $0 \leq b$ (at%) < 5 .

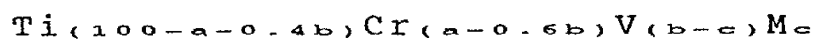
4. The hydrogen storage metal alloy according to Claim 3 wherein a level each of the constituent element Mo and/or W contained in the alloy is within a range of 3 ± 1.5 at %.

5. A hydrogen storage metal alloy which has
(1) as its main phase a body-centered cubic structure-type phase enabling the adsorption and desorption of hydrogen, and

30

(2) a composition of the following general composition formula:

35



wherein M is at least a member selected from molybdenum (Mo) and tungsten (W), provided that $20 \leq a$ (at%) ≤ 80 , $0 \leq b$ (at%) ≤ 10 , and $0 \leq c$ (at%) < 5 .

5 6. The hydrogen storage metal alloy according to any of Claims 1 to 5 wherein the element X is admixed at its atom % concentration, d (at%), ranging within $0 \leq d$ (at%) ≤ 20 , the atomic radius of which is larger than that of Cr but smaller than that of Ti.

10

 7. The hydrogen storage metal alloy according to Claim 6 wherein the element X is at least one or more members selected from the group consisting of Al, Ge, Ga, Si, Au and Pt.

15

 8. The hydrogen storage metal alloy according to any of Claims 1 to 7 wherein the element T is admixed at its atom % concentration, e (at%), ranging within $0 \leq e$ (at%) ≤ 10 and includes at least one or
20 more members selected from the group consisting of Nb, Ta, Mn, Fe, Al, B, C, Co, Cu, Ga, Ge, Ln (a variety of lanthanoid metals), N, Ni, P, and Si.

20

 9. A process for producing a hydrogen storage
25 metal alloy which is for the production of the hydrogen storage metal alloy having as the main phase a body-centered cubic structure-type phase enabling the adsorption and desorption of hydrogen, and

 comprises the steps of:

30

 (1) melting a starting alloy brought to a predetermined element ratio to form a uniform heat (melting step),

35

 (2) keeping the homogenized alloy heat at a temperature within a range just below the melting point of the alloy for a predetermined time (heat treatment), and

 (3) rapidly cooling the alloy after the heat

treatment (quenching step).

10. The process of Claim 9 wherein melting and solidification may be carried out repeatedly
5 predetermined times at the aforementioned melting step.

11. The process of Claim 9 or 10 wherein the predetermined time range at the aforementioned heat treatment is from 1 minute to 100 hours.

12. The process according to any of Claims 9 to 11 wherein the element ratios are those described in any of Claims 1 to 8 regarding the hydrogen storage metal alloy products.

A process for producing hydrogen storage metal alloys having a body-centered cubic structure-type main phase enabling the adsorption and desorption of hydrogen is provided which comprises the steps of: (1) melting a starting alloy brought to a predetermined element ratio to form a uniform heat (melting step), (2) keeping the homogenized alloy heat at a temperature within a range just below the melting point of the alloy for a predetermined time (heat treatment), and (3) rapidly cooling the alloy after the heat treatment (quenching step).

See: FIG. 1

Figure 1. The effect of the number of trials on the number of correct responses. The number of correct responses was plotted against the number of trials for each condition. The number of correct responses increased with the number of trials for all conditions. The number of correct responses was highest for the condition with the highest number of trials (10 trials) and lowest for the condition with the lowest number of trials (2 trials).

FIG. 1

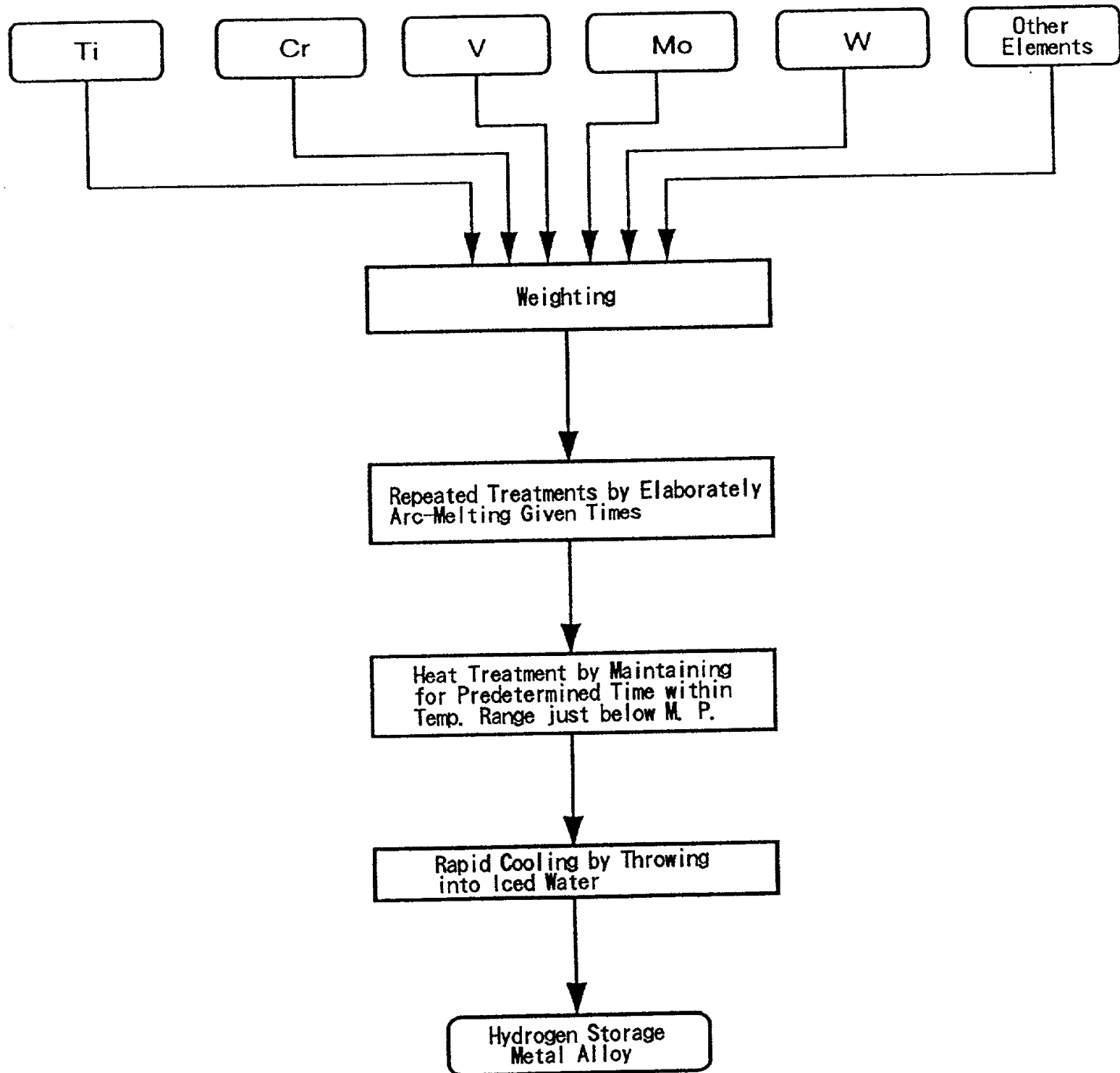


FIG. 2

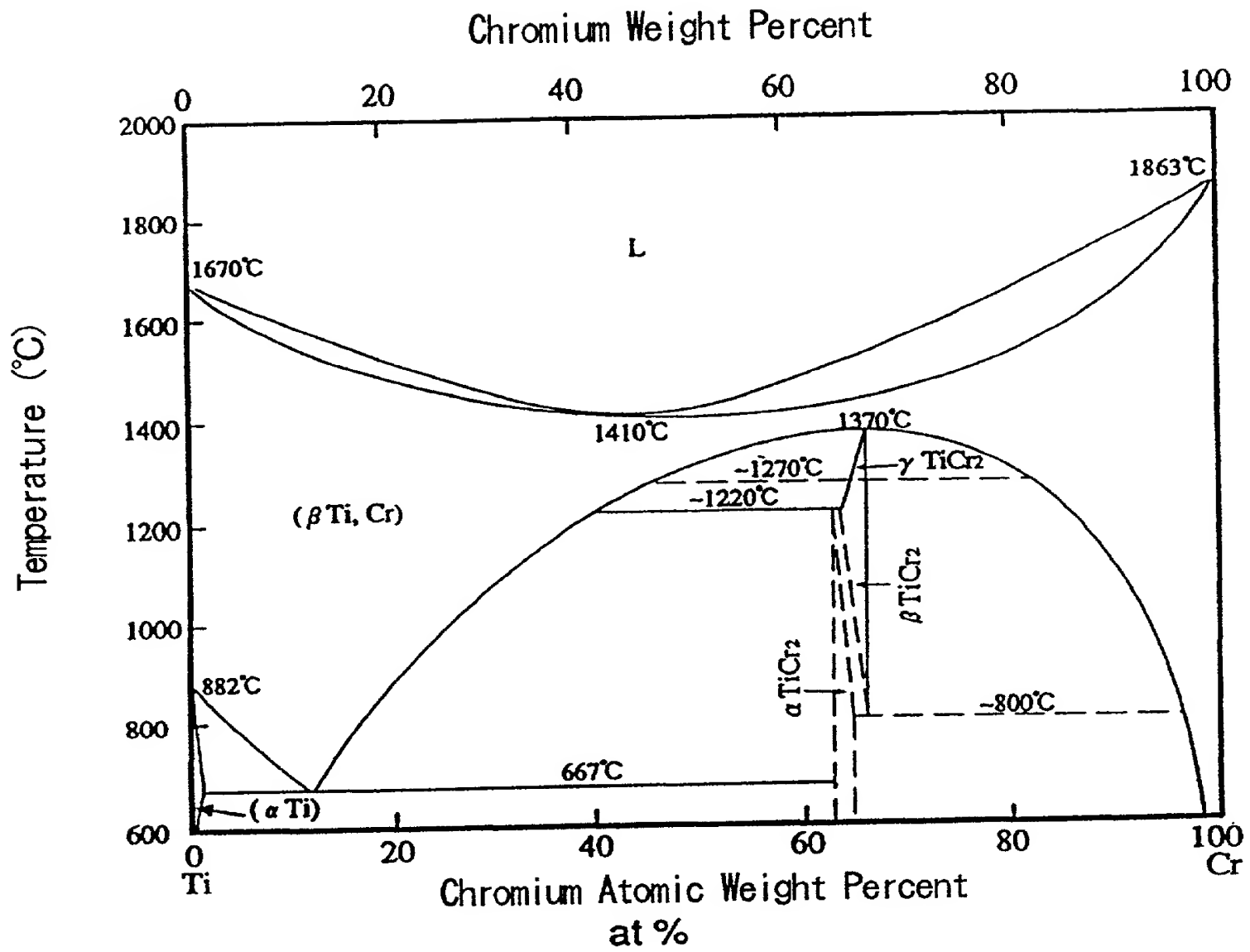


FIG. 3

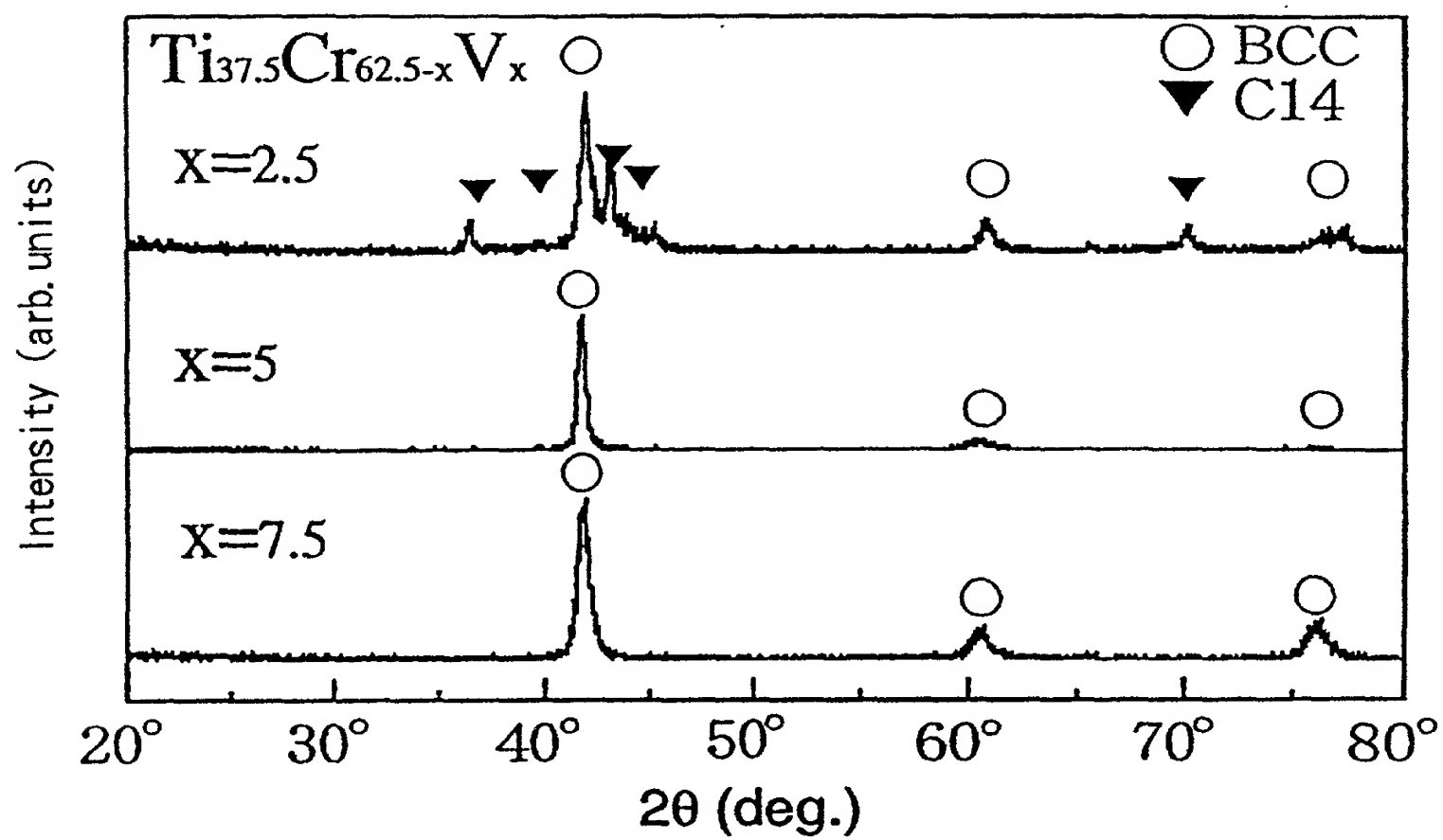


FIG. 4

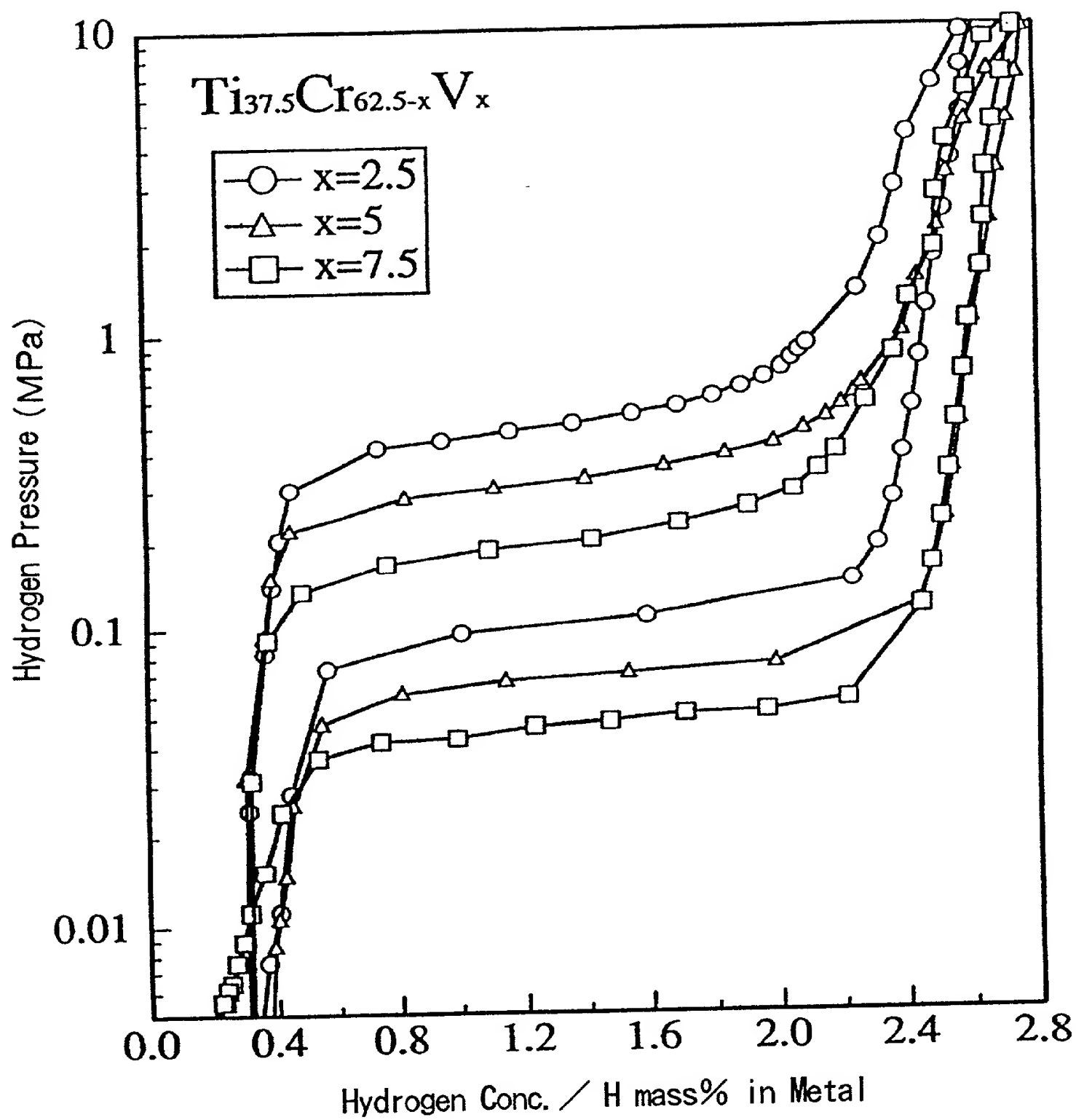


FIG. 5

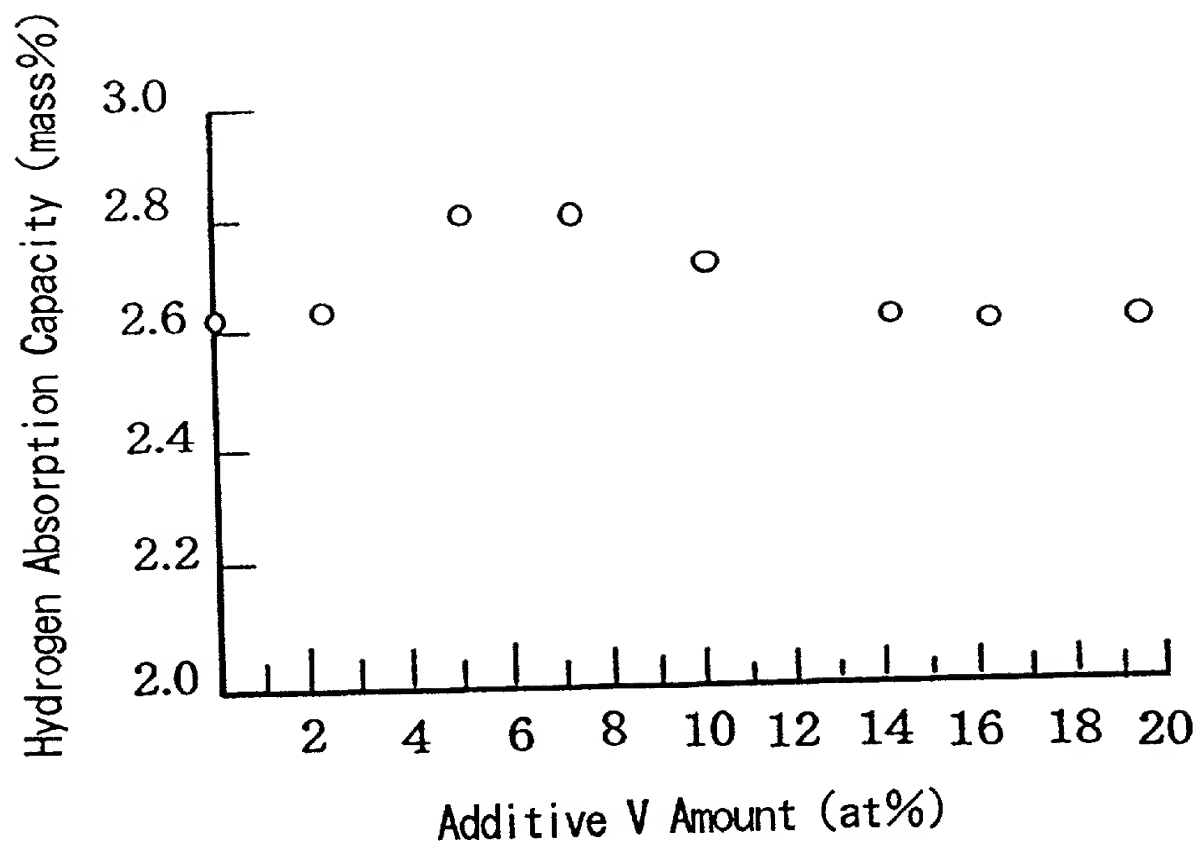


FIG. 6

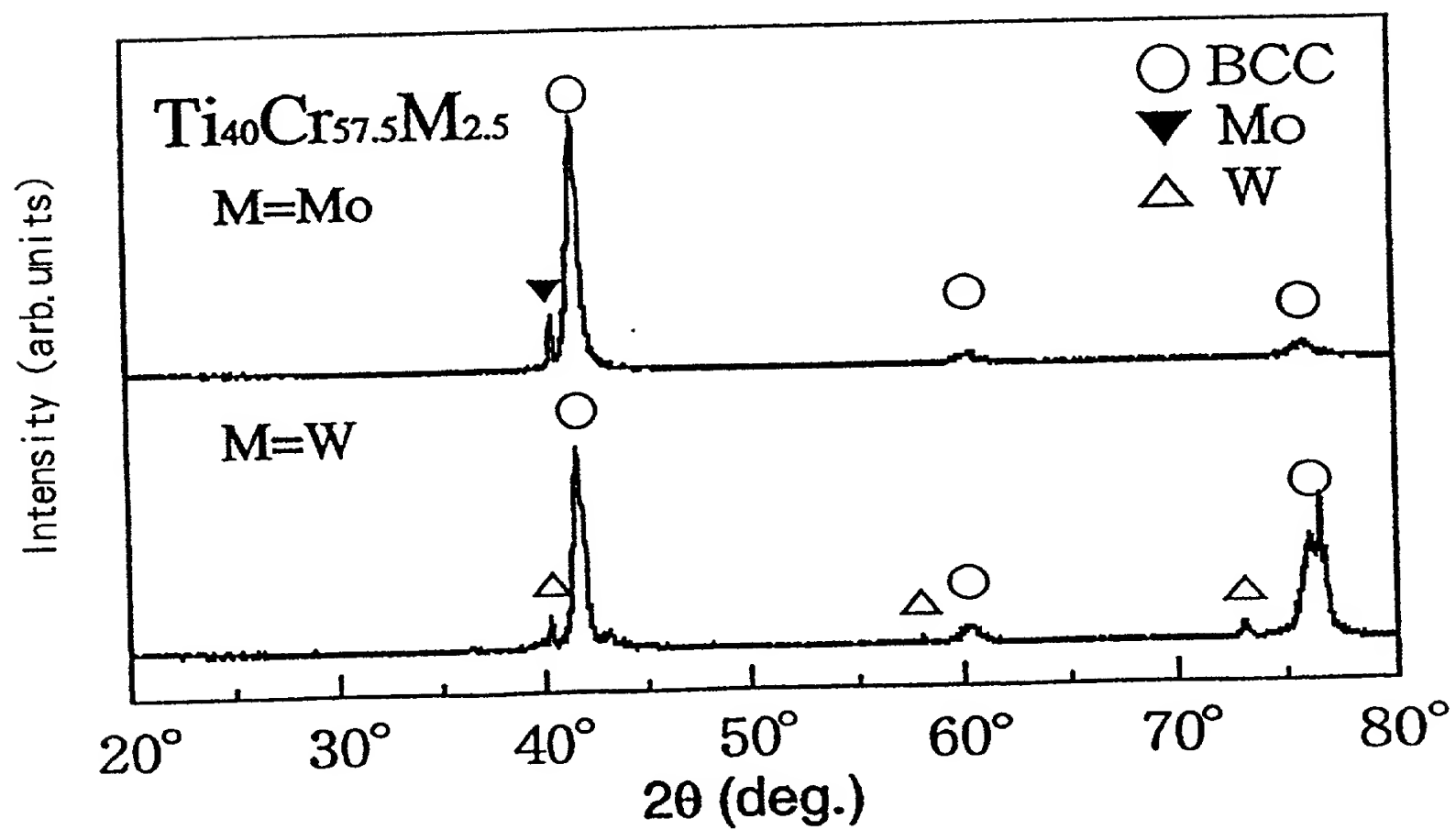


FIG. 7

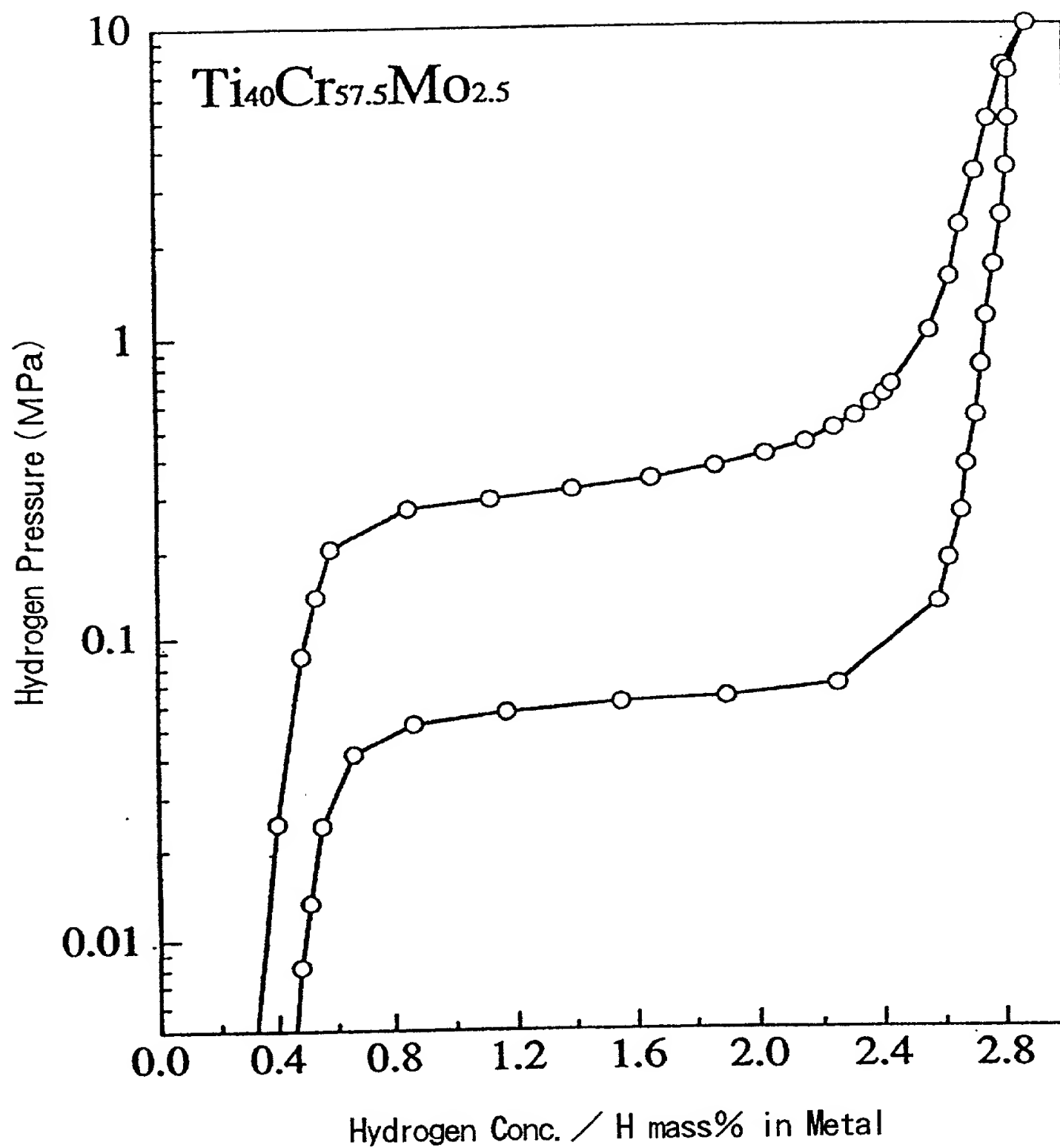


FIG. 8

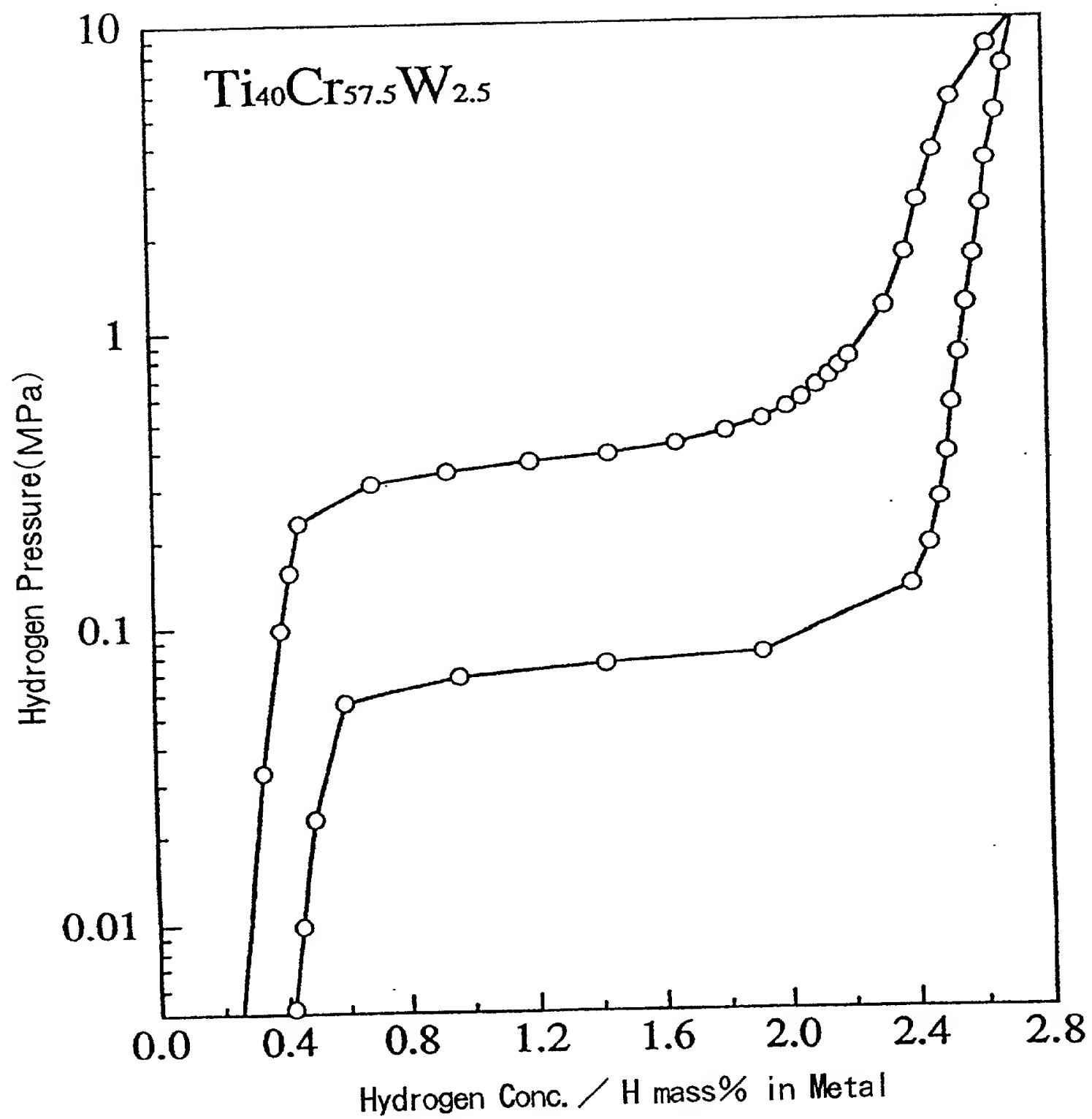


FIG. 9

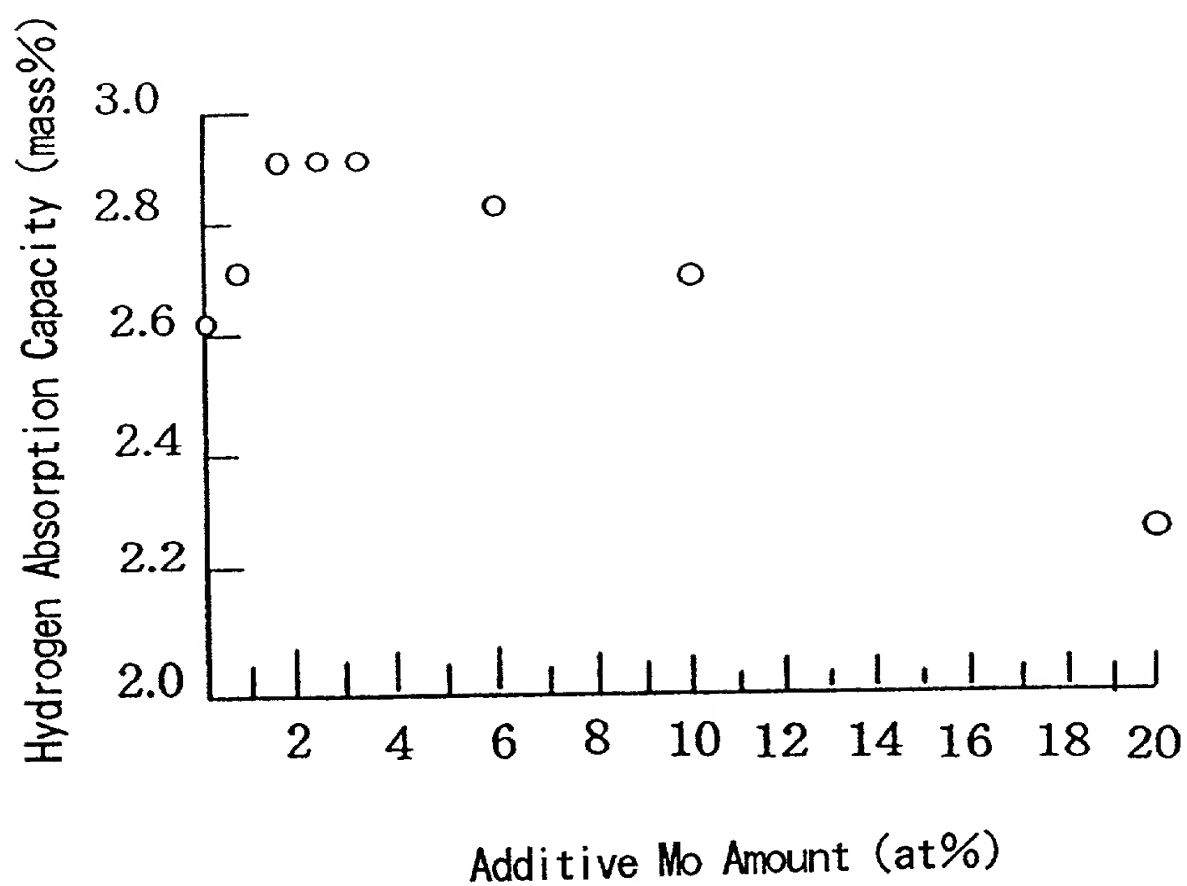


FIG. 10

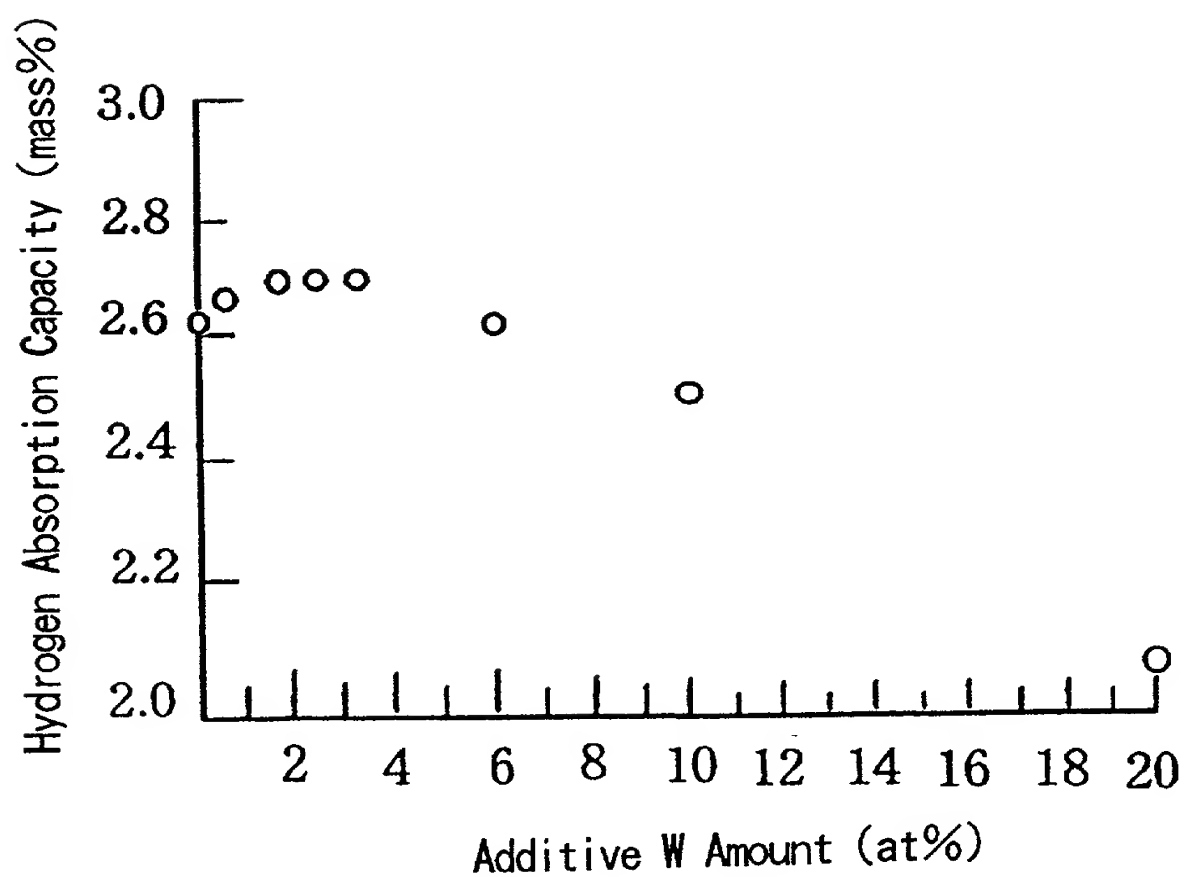


FIG. 11

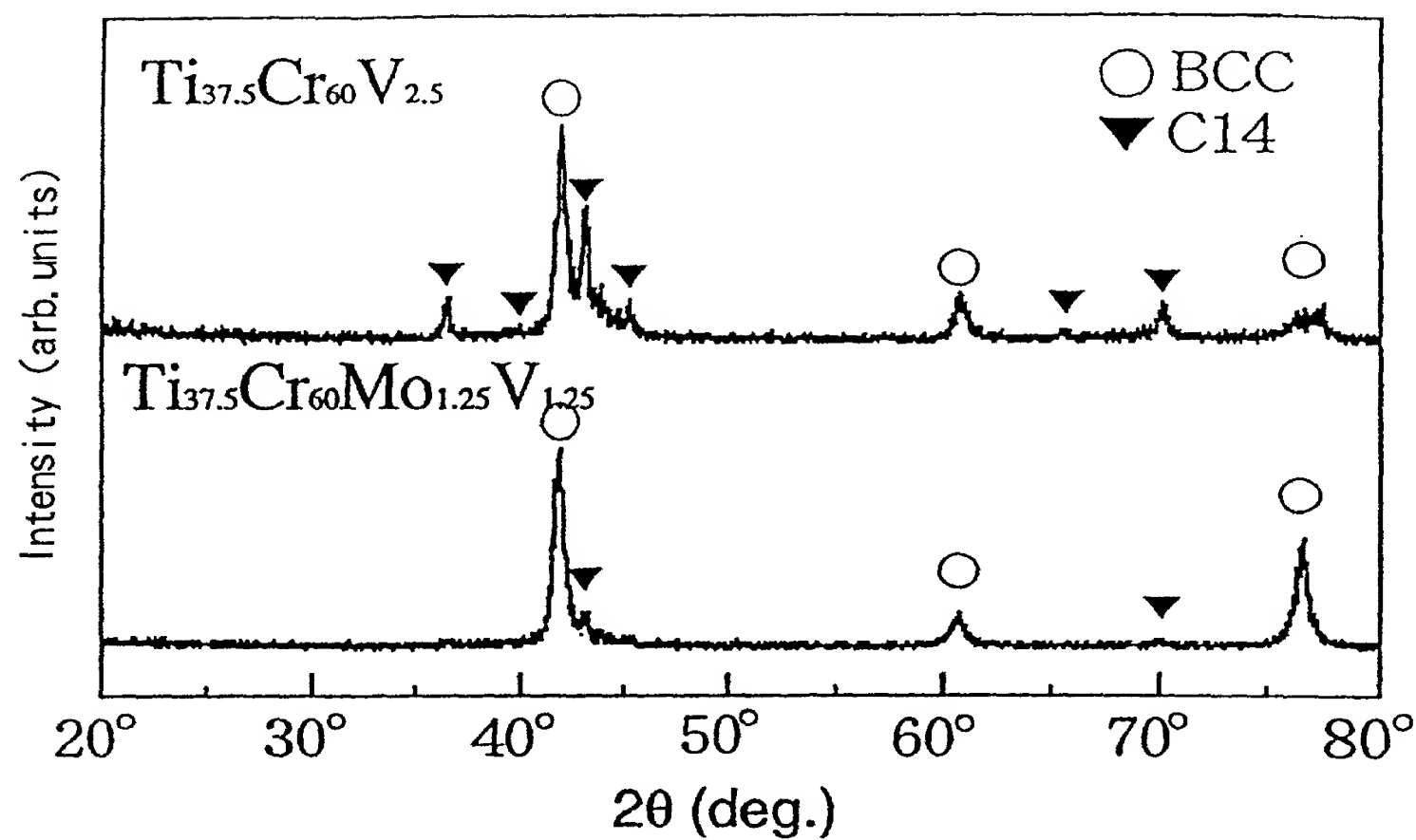


FIG. 12

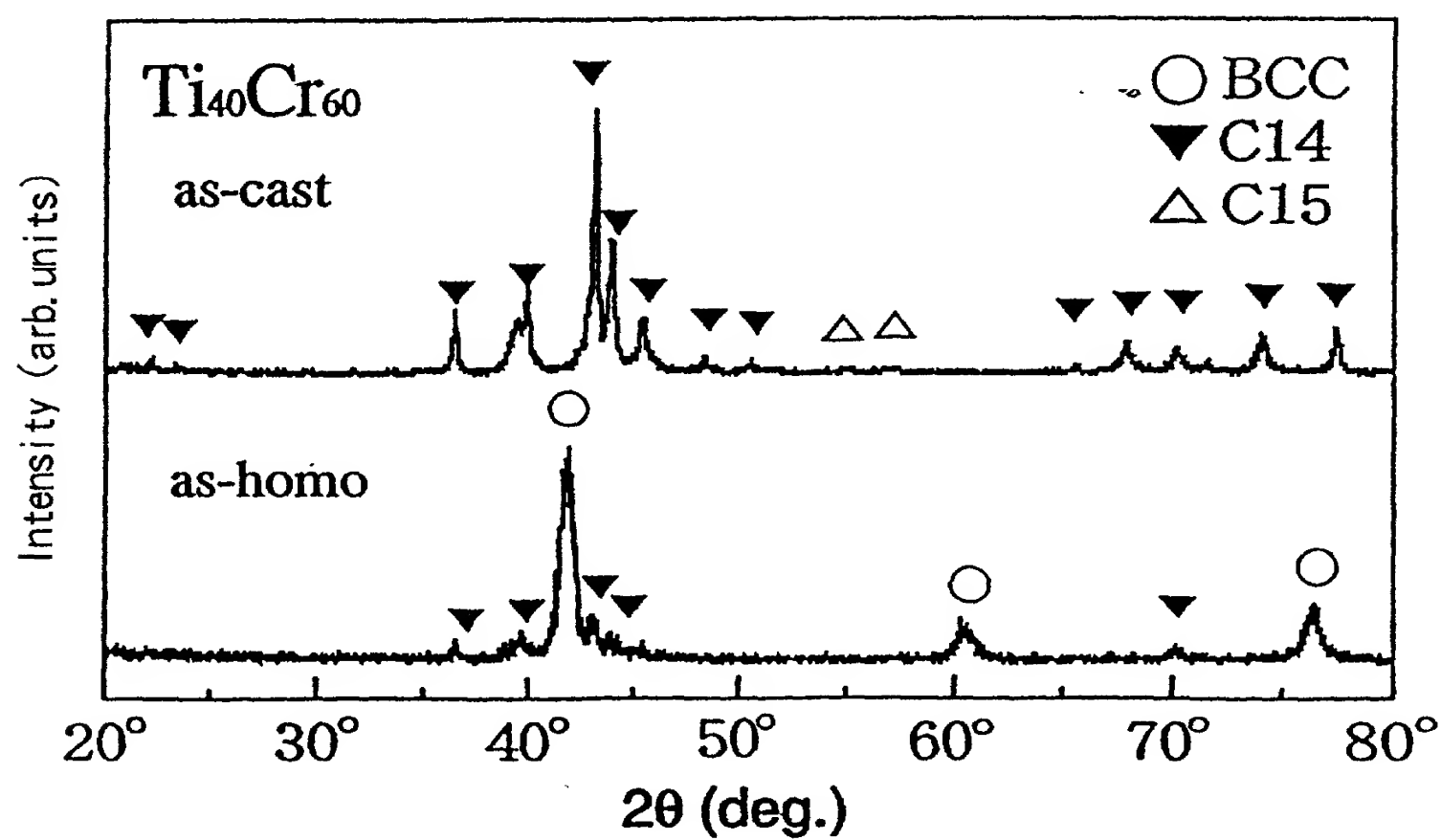


FIG. 13

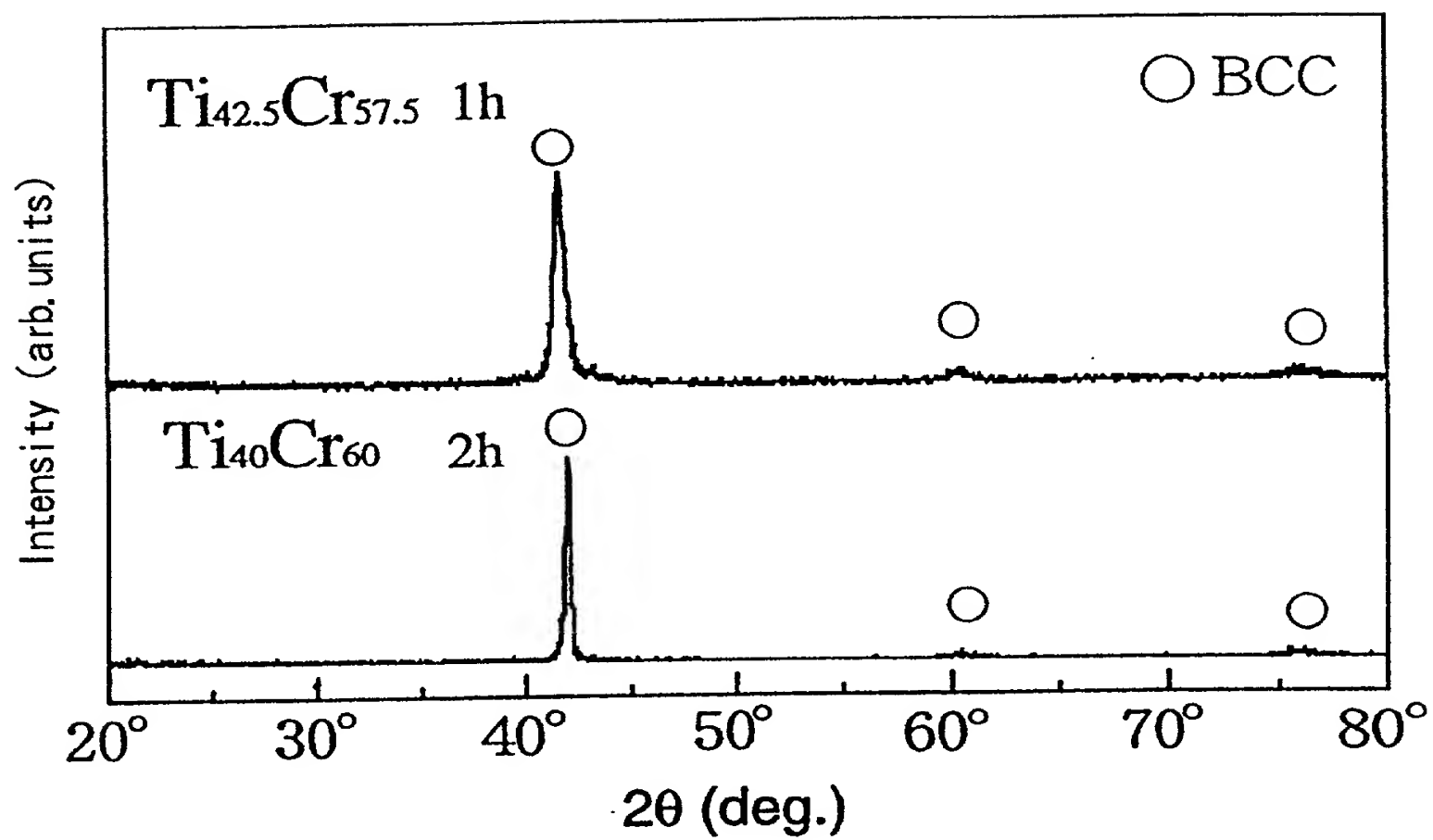


FIG. 14

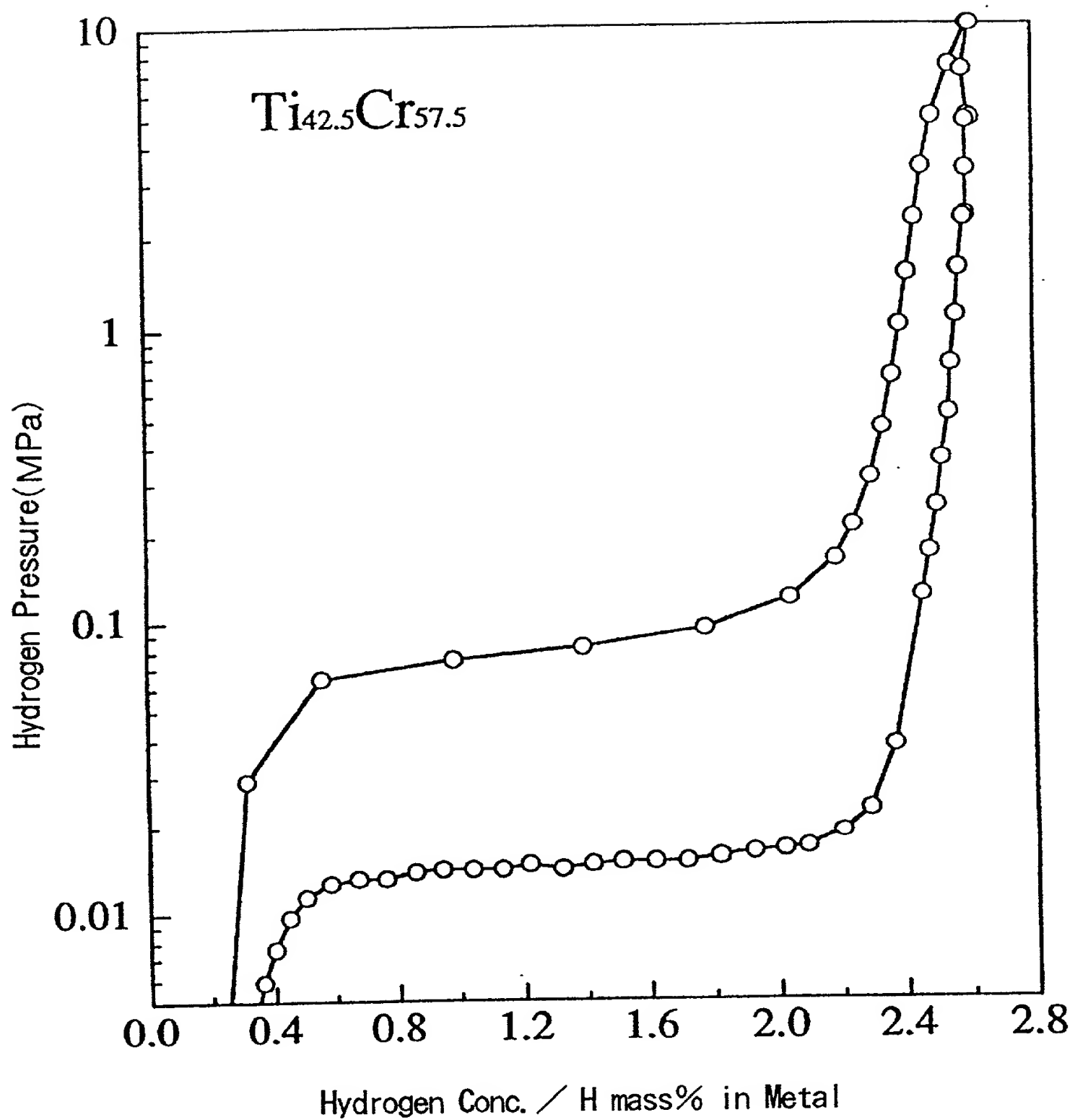


FIG. 15

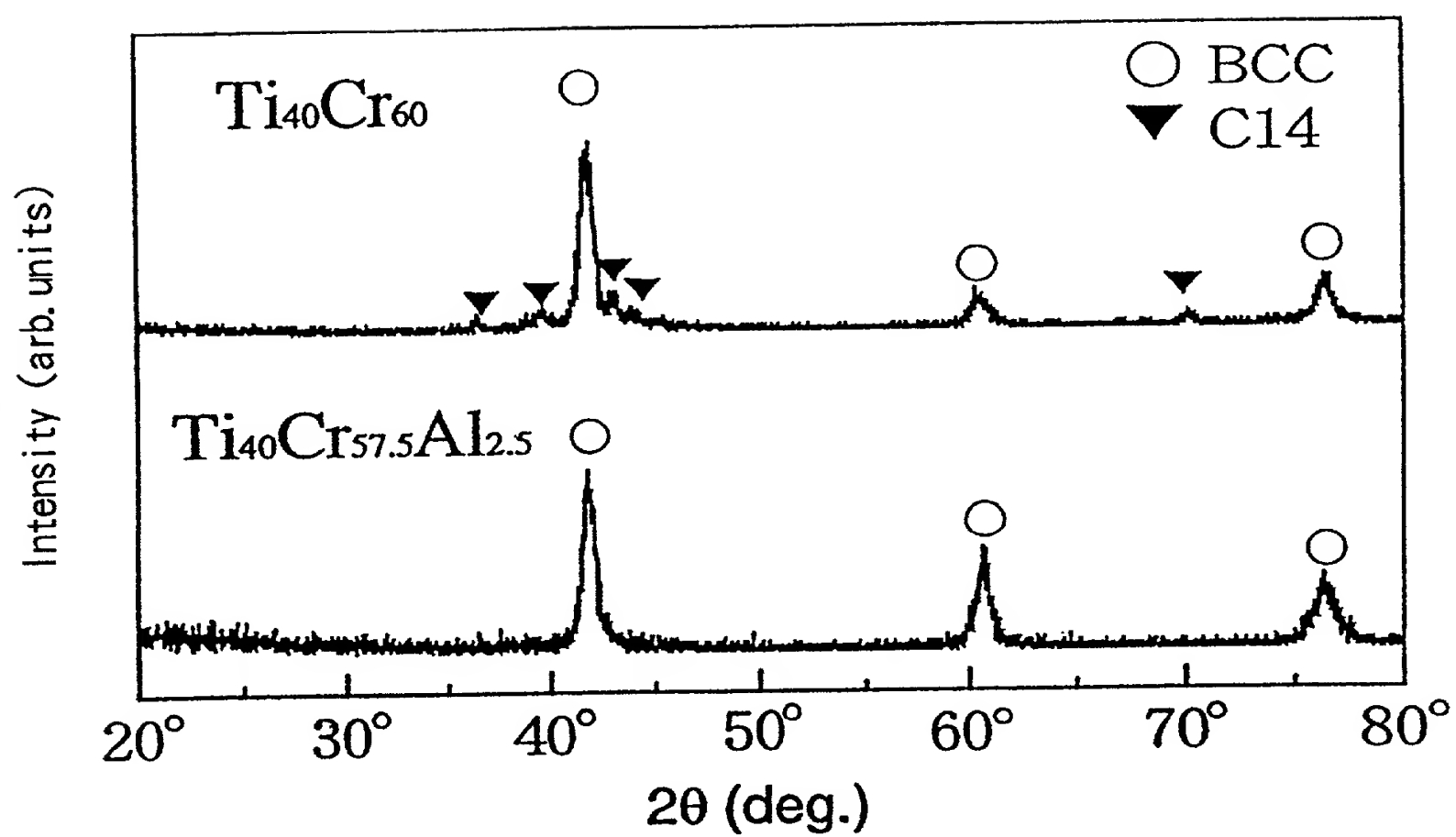
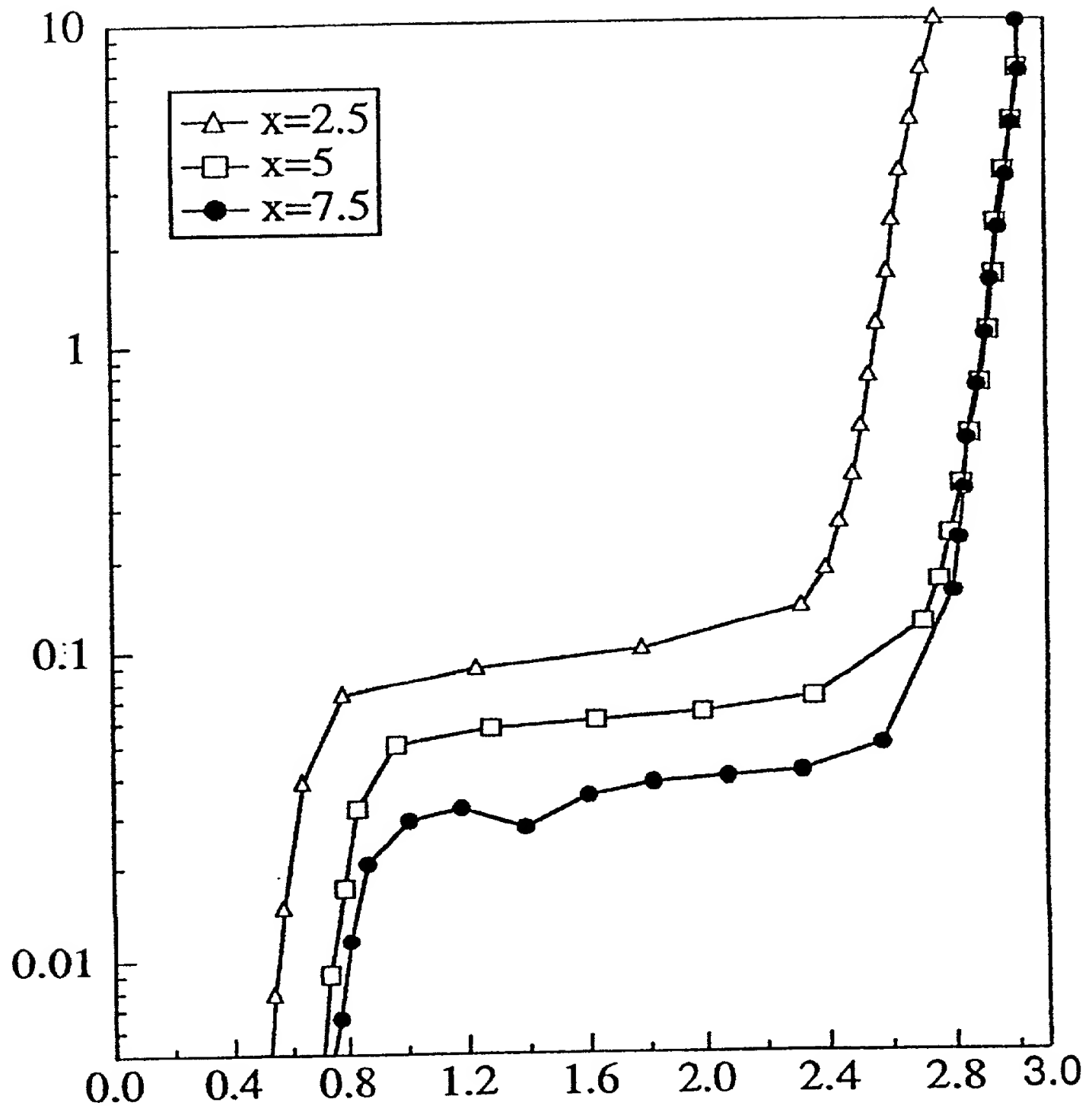


FIG. 16



DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION

Attorney Docket No: SHIG C11119

First Named Inventor: OKADA et al

Complete if known: Serial No: _____ Filing Date: August 16, 2001

Group Art Unit: _____ Examiner: _____

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name.

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled **HYDROGEN STORAGE METAL ALLOY AND PRODUCTION THEREOF**, the specification of which is attached hereto.

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, S. 1.56(a).

I hereby claim foreign priority benefits under 35 U.S.C. 119(a)-(d) or 365(b) of any foreign application(s) for patent or inventor's certificate, or 365(a) of any PCT international application which designated at least one country other than the United States of America, listed below and have also identified below any foreign application for patent or inventor's certificate or of any PCT international application having a filing date before that of the application on which priority is claimed:

Prior Foreign Application(s):

<u>Prior Foreign Application(s):</u>			<u>Priority Claimed</u>	<u>Certified Copy Attached</u>
<u>11-359899</u>	<u>JAPAN</u>	<u>12/17/1999</u>	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No	<input type="checkbox"/> Yes <input type="checkbox"/> No
(Number)	(Country)	(Month/Day/Year Filed)		
_____	_____	_____	<input type="checkbox"/> Yes <input type="checkbox"/> No	<input type="checkbox"/> Yes <input type="checkbox"/> No
(Number)	(Country)	(Month/Day/Year Filed)		

I hereby claim the benefit under 35 U.S.C. 119(e) of any United States provisional application(s) listed below:

Application No:

Filing Date:

I hereby claim the benefit under 35 U.S.C. 120 of any United States application(s), or 365(c) of any PCT international application designating the United States of America, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of 35 U.S.C. 112, I acknowledge the duty to disclose information which is material to patentability as defined in 37 CFR 1.56 which became available between the filing date of the prior application and the national or PCT international filing date of this application:

PCT/JP00/02056

03/30/2000

US Parent Application No.
or PCT Parent Appln. No.

Parent Filing Date

Parent Patent Number
(if applicable)

And I hereby appoint HAYES, SOLOWAY, HENNESSEY, GROSSMAN & HAGE, P.C., a firm composed of Oliver W. Hayes, Reg. No. 15,867; Norman P. Soloway, Reg. No. 24,315; William O. Hennessey, Reg. No. 32,032; Susan H. Hage, Reg. No. 29,646; Steven J. Grossman, Reg. No. 35,001; and Donald J. Perreault, Reg. No. 40,126, or any of them, of 175 Canal Street, Manchester, New Hampshire 03101 (Telephone: 603-668-1400); or Edmund Paul Pfleger, Reg. No. 41,252; Dale F. Regelman, Reg. No. 45,625; or Kevin M. Drucker, Reg. No. 47,537, or any of them, of 130 W. Cushing Street, Tucson, Arizona 85701 (Telephone: 520-882-7623) my attorneys with full power of substitution and revocation, to prosecute this application and to transact all business in the Patent Office connected therewith.

Please direct all future correspondence in connection with this application to the attention of Norman P. Soloway, HAYES, SOLOWAY, HENNESSEY, GROSSMAN & HAGE, P.C., 130 W. Cushing Street, Tucson, Arizona 85701 (Telephone: 520-882-7623).

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

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FD-973 (Rev. 10-1-95)

IMPORTANT NOTICE RE DUTY OF CANDOR AND GOOD FAITH

The Duty of Disclosure requirements of Section 1.56(a), of Title 37 of the Code of Federal Regulations are as follows:

A duty of candor and good faith toward the Patent and Trademark Office rests on the inventor, on each attorney or agent who prepares or prosecutes the application and on every other individual who is substantively involved in the preparation or prosecution of the application and who is associated with the inventor, with the assignee or with anyone to whom there is an obligation to assign the application. All such individuals have a duty to disclose to the Office information they are aware of which is material to the examination of the application. Such information is material where there is a substantial likelihood that a reasonable examiner would consider it important in deciding whether to allow the application to issue as a patent. The duty is commensurate with the degree of involvement in the preparation or prosecution of the application.

By virtue of this regulation each inventor executing the Declaration for the filing of a Patent Application acknowledges his duty to disclose information of which he is aware and which may be material to the examination of the application.

Inherent in this is the duty to disclose any knowledge or belief that the invention:

- (a) was ever known or used in the United States of America before his invention thereof;
- (b) was patented or described in any printed publication in any country before his invention thereof or more than one year prior to the actual filing date of the U.S. patent application;
- (c) was in public use or on sale in the United States of America more than one year prior to the actual filing date of the U.S. patent application; or
- (d) has been patented or made the subject of inventor's certificate issued before the actual filing date of the U.S. patent application in any country foreign to the United States of America on an application filed by him or his legal representatives or assigns more than twelve months before the actual filing date in the United States.

NOTE: The "Information" concerned includes, but is not limited to, all published applications and patents, including applicant's and assignee's own U.S. or foreign applications and patents, as well as any other pertinent prior art known, or which becomes known, to the inventor or his representatives. Where English language equivalents of foreign language documents are known, they should be identified and, when possible, copies supplied. Failure to comply with this requirement may result in a patent issued on the application being held invalid even if the known prior art which is not supplied is material to only one claim of that patent.

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